Real-time intracellular temperature imaging using lanthanide-bearing polymeric micelles

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Measurement of thermogenesis in individual cells is a remarkable challenge due to the complexity of the biochemical environment (such as pH and ionic strength) and to the rapid and yet not well-understood heat transfer mechanisms throughout the cell. Here, we present a unique system for intracellular temperature mapping in a fluorescence microscope (uncertainty of 0.2 K) using rationally designed luminescent Ln³+-bearing polymeric micellar probes (Ln=Sm, Eu) incubated in breast cancer MDA-MB468 cells. 2D thermal images recorded increasing the temperature of the cells culture medium between 296 and 304 K shows inhomogeneous intracellular temperature progressions up to ~20 degrees and subcellular gradients of ~5 degrees between the nucleolus and the rest of the cell, illustrating the thermogenic activity of the different organelles and highlighting the potential of this tool to study intracellular processes.

Keywords: Intracellular temperature mapping, Luminescence thermometry, Thermogenesis, Lanthanide-based polymer nanomicelles

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Temperature affects everything and temperature fluctuations play a central role in the myriad of biochemical reactions regulating life.¹ The intracellular temperature, for instance, changes with cellular activity, including cell division, gene expression, enzymatic reactions, and pathological states.²-⁴ Moreover, to neutralize large external temperature changes and to keep homeostasis of the body temperature upon cold or hot exposure, cells have developed thermoregulation mechanisms.³-7 Despite recognition of the fundamental involvement of temperature in eliciting biochemical changes,³-10 specific molecular mechanisms by which cells produce and use heat are largely unknown,⁴. 7, ¹¹¹-¹³, which, thus, have driven the research on intracellular thermometers in the past decade. Although there are a handful of reported intrusive intracellular thermometers,⁶. ¹³, ²0 contactless luminescence-based examples have been largely the most adopted strategy.³, ¹¹0, ²¹-²7 Indeed, several types of luminescent materials have been proposed for intracellular temperature sensing such as organic molecules,²¹, ²³, ²³ polymers,³₀, ³¹ carbon dots,³² quantum dots,³³ N-vacancy nanodiamonds,² and trivalent lanthanide ions (Ln³+)-doped nanomaterials.³⁴

These luminescent thermometers unveiled intriguing spatiotemporal intracellular temperature fluctuations, including organelle-specific thermogenesis, suggesting a fundamental connection between the temperature and the cell functions.^{4, 7-10} For example, Chrétien *et al.*^{4, 7-10} correlated changes in the mitochondrial temperature with the oxygen consumption, showing that the mitochondrial temperature is up to 10 degrees higher than the body temperature when the respiratory chain was fully functional. Also, Tanimoto *et al.*³⁵ observed temperature differences up to 6 degrees in micrometer scale domains in a neuronal cell after the mitochondrial stimulation, whereas Rajagopal *et al.*⁶ reported temperature spikes of ~7.5 degrees during stimulated proton uncoupling in neurons of *Aplysia californica*. Finally, Yang *et al.*³³ and Liu *et al.*³⁶ showed inhomogeneous intracellular temperature distributions up to 10–15 degrees higher than the temperature of the living cells culture medium, following, respectively, Ca²⁺ stress and cold shock, and externally-induced temperature changes.

All these observations (among other examples^{5, 30}) are inconsistent with theoretical calculations suggesting that maximal heat gradients across cells could not be greater than 10^{-5} K.^{5, 30}

Here, we report intriguing real-time bidimensional (2D) temperature maps of breast metastatic adenocarcinoma cells (line MDA-MB-468) incubated with innovative Ln³⁺-bearing polymeric micelles. The thermal images show inhomogeneous intracellular temperature progressions up to ~20 degrees and subcellular temperature gradients of ~5 degrees raising the exciting possibility of using these gradients for signaling and regulation in cells,³³ and emphasizing the present misunderstood about the response of living cells to temperature variations.

A scheme of the synthesis of the micelles is presented in Figure 1. It consists of a self-organized structure formed using a single polymer containing hydrophilic and hydrophobic blocks. The amphiphilic block is the P(MPEGA-co-PEGA)-b-P(MA-b-PhenA) copolymer (MPEGA and PEGA stand for poly(ethylene glycol) methyl ether acrylate and polyethyleneglycol acrylate, respectively) and was synthesized by reversible addition-fragmentation chain transfer polymerization (details in the Supporting Information). The hydrophobic block is a methyl acrylate (MA) polymer containing a discrete number of the auxiliary ligand 4-(acryloyloxymethyl)-1,10phenanthroline (PhenA) to complete the coordination sphere of the Ln3+ ion and shield the complex from the influence of the medium by steric effects. We rationally design the polymeric micelles and the Ln3+ complexes to mitigate luminescence quenching effects induced by the culture medium and the proposed copolymerization strategy permits easy incorporation of other functionalities such as tags for different cell organelles. The hydrodynamic diameter of the resulting micelles in aqueous suspension gives a monomodal distribution with an average diameter of 20.3±0.2 nm (Figure 1b,c). The scanning transmission electron micrographs showed particles, with a granular texture and bright spots (discernible at high magnification), which is compatible with the presence of Ln3+ complexes within the micelle. Moreover, the electron dispersive scattering (EDS) within the micelles presents the signature of Sm, whereas less frequent Eu, was only faintly detected (Figure S2 in Supporting Information). The experimental ratios of the Ln³⁺ ions were 2.42% and 2.84% w/w, for DNDP and btfa complexes, respectively. These elements were not observed outside the micelles indicating that there is no leakage of the Ln³⁺ ions to the suspension (Figures S3-S5 in Supporting Information).

The photoluminescence studies on the Ln³+-bearing polymeric micelles are summarized in Figure 2. A remarkable excitation peak at 436 nm, in the blue-violet spectral range, is discerned (Figure 2a). This is an important achievement because contrary to other down-shifting Ln³+-based thermometers that need high-energetic (and cell harmful) ultraviolet excitation, ³⁷ the excitation in the visible spectral range circumvents damage to the cultured cells under observation and it can be easily used in conventional confocal microscopes and cytometers that do not natively incorporate UV laser sources. Moreover, as these instruments do not also generally incorporate NIR laser sources, intracellular thermometers based on Ln³+-doped upconverting nanoparticles³4 and nanocomposites³8-40 (generally excited at 980 nm) cannot be used routinely, despite the negligible autofluorescence of the cells induced by NIR excitation.

The thermal dependence of the emission spectra was recorded in the 300–328 K range for the aqueous suspension of the Eu³⁺- and Sm³⁺-bearing polymeric micelles (Figure 2b-d). In the case of btfa-based micelles, the integrated area of the Sm³⁺ ${}^4G_{5/2} \rightarrow {}^6H_{9/2}$ transition (I_1) decreases with increasing temperature by ~12%, that of the Eu³⁺, ${}^5D_0 \rightarrow {}^7F_2$ transition (I_2) is reduced by 40% of the initial value (Figure 3a). According to the DNPD-based material, the temperature-dependent trend is more evident. The emission of Eu³⁺ decreases increasing temperature (leaving ~18% of the RT intensity), and what is more also the emission of Sm³⁺ is reduced ~4 times at the highest temperature (Figure 3b). To rationalize these thermal dependences, we use time-dependent density functional theory and intramolecular energy transfer

calculations (details in Supporting Information). The intramolecular energy transfer rates for Ln³+ complexes were estimated based on a well-defined theory.⁴¹-⁴⁵

The main energy-transfer pathways in the micelles containing the DNPD complexes (Figure 3c) involves the $T_1 \rightarrow {}^5D_0$ (Eu³⁺) and $T_1 \rightarrow {}^4G_{5/2}$ (Sm³⁺) transitions corresponding respectively to ~83% and ~91% of the total forward intramolecular energy transfer rates. On one hand, the backward energy transfer rates (W_{bET}) increase at temperature increase, depopulating the ⁵D₀ (Eu³⁺) and ⁴G_{5/2} (Sm³⁺) emitting levels and, on the other hand, $W_{\rm bET}$ for Eu³⁺ is more affected by the temperature increase than those of Sm3+. The balance between these two processes explains the stepper decay observed for the integrated area of ${}^5D_0 \rightarrow {}^7F_2$ transition in comparison with that of ${}^4G_{5/2} \rightarrow {}^6H_{9/2}$ (Figure 3b). In the btfa-Ln³⁺ complexes, the T₁ state (15480 cm⁻¹) is lying below the Eu³⁺ and Sm³⁺ ion acceptor states, leading to a higher energy barrier. Consequently, the energy transfer pathways involving the S₁ state (34300 cm⁻¹) as donor become more operative (in order of 10⁴ and 10⁶ s⁻¹, respectively, Tables S5 and S6). According to the simulations, the energy transfer from the S₁ represents 93.4% and 93.8% of the total for Eu³⁺ and Sm³⁺ ions, respectively. For the btfa ligand, the energy transfer involves the non-radiative decays from the lanthanides' higher energy states to the corresponding emitting levels. Since the population of the emitting levels is dependent on its decay lifetimes (that are in order of 10⁻³ s), it has time enough to transfer energy to the triplet state (in order of 10⁴ s⁻¹ for Eu³⁺ and 10⁵ s⁻¹ for Sm³⁺). This means that the population of the emitting level is dependent on the balance between the population increase resulting from the singlet (Win) and the population decrease due to lanthanide-to-btfa triplet state ($W_{T_1}^{Ln}$) energy transfers, which allow us to define an effective transition rate as $W^{\ast}=W_{Ln}^{S}-W_{T_{1}}^{Ln}\,.$ Figure S7 presents the temperature dependence of the normalized W^* . The rate for the $Eu^{3+} \rightarrow T_1$ energy transfer is of the same order of magnitude of that of $S_1 \rightarrow Eu^{3+}$ in the higher energetic states. In contrast, for the btfa-Sm $^{3+}$ complex, the $W^{Ln}_{T_1}$ is one order of magnitude lower than the W_{Ln}^S , explaining the lower temperature quenching of the Sm³⁺ integrated area in comparison with the Eu³⁺ one presented in Figure 3a.

The integrated areas of the ${}^4G_{5/2} \rightarrow {}^6H_{9/2}$ (I_1 , Sm³+) and ${}^5D_0 \rightarrow {}^7F_2$ (I_2 , Eu³+) transitions are used to define the thermometric parameter $\Delta = I_1/I_2$. The Δ (T) dependence is well fitted by empirical polynomial equations for both ligands (Figure S13 in Supporting Information). The relative thermal sensitivity (Figure S14 in Supporting Information) of the micelles increases with the temperature increase reaching the maximum value of 1.5 %. K⁻¹ and 1.7 %. K⁻¹ for btfa- and DNPD-containing micelles, respectively, both at 328 K. The minimum temperature uncertainty is remarkably low, 6 mK (at 328 K), and 0.20 K (at 305 K) for the btfa- and DNPD-containing micelles (Figure S16 in Supporting Information). Whereas the relative thermal sensitivity values are lower than that reported for similar micelles containing Tb³+ ions³7 (5.8 %·K⁻¹ at 296 K), the decrease of the relative uncertainty on the integrated intensity to unprecedented values of $\delta I/I < 0.4\%$ (Figure S15 in Supporting Information) allowed us to design the first luminescent thermometer with temperature uncertainty values below 10 mK.

These exciting results of thermometric performance motivate the development of a temperature-imaging instrument (Figure S7 in Supporting Information). It consists of a fluorescence microscope that incorporates a beam splitter, a CMOS camera, and a CCD camera. The illumination light is filtered through a 340–380 nm bandpass filter and reflected with a 400 nm dichroic mirror. The emission beam is divided into two by a 635 nm dichroic mirror and the resulting beams are filtered through the 600–620 nm and 620–660 nm bandpass filters that roughly cover the $^5D_0 \rightarrow ^7F_2$ (Eu³⁺) and $^4G_{5/2} \rightarrow ^6H_{9/2}$ (Sm³⁺) emissions. Finally, both the Eu³⁺ and Sm³⁺ filtered beams are collected by a" CMOS camera (Figure S7b in Supporting Information). All the process of temperature imaging is controlled by a LabView image-processing sequence. A MatLab® routine (developed by us) is then used to process the original CMOS image containing both the Eu³⁺and Sm³⁺ emission to get 2D temperature maps (Figure S7b in Supporting Information). The temperature-mapping routine first separates the original

image into the corresponding Sm³⁺ (I_1) and Eu³⁺ (I_2) emissions. Then, the intensity values for the equivalent pixels of Eu³⁺ and Sm³⁺ images are identified, and the intensity ratio $\Delta = I_1/I_2$ is calculated and converted into temperature by the corresponding calibration curve. Finally, we obtain a 2D temperature matrix that is represented as a false-color image that can be overlapped to the original image, mapping the temperature distribution of the cellular culture.

Despite both the btfa- and DNPD-based Ln³+-bearing polymeric micelles are stable in water, in the presence of the culture medium the btfa-based polymeric micelles present a poor emission intensity in the same excitation conditions of DNPD-based ones (Figure S22 in Supporting Information). The time-dependent density functional theory and the intramolecular energy transfer calculations point out that the btfa-based micelles undergo a luminescence quenching process due to the energy transfer that involves the Ln³+ emitting levels to the T¹ state. On the contrary, in the DNPD-based micelles is the balance between the S¹-to-Ln³+ and the Ln³+-to-T¹ energy transfers that rules the temperature dependence of the intensities (details *in silico* experiments in Supporting Information).

In a proof-of-concept experiment, the thermometric imaging system was tested with MDA-MB-468 breast cancer cells incubated with DNPD-based polymeric micelles. The cells were observed before and after incubation with the Ln³+-bearing polymeric micelles (culture and cell internalization details in Supporting Information). Phase-contrast images (Figure 4a,b) showed that the cells were well-formed and uniformly spread. Grayscale images showed the luminescence of internalized Ln³+-bearing polymeric micelles in both the Sm³+ and the Eu³+ channels (Figure 4c,d, and Figure S23c,d in Supporting Information), whereas images from control cells before incubation with the micelles were totally dark (Figure S18 in Supporting Information). These results are compatible with the cell internalization of the Ln³+-bearing polymeric micelles and indicate their stability on cellular medium. Moreover, the mean temperature determined in both the fixed and live cells agrees very well with the

temperature values of the cellular culture (298.0 and 296.5 K, respectively, Figures S19, S23 and S24 in Supporting Information). Interestingly, the temperature distribution is much narrower in the fixed cells (standard error of 0.02 K, Figures S19 and S23 in Supporting Information) than in the live cell (standard error >0.5 K Figure 4c,d and Figure S24 in Supporting Information).

The phase-contrast images of the cell cultures incubated with DNPD-based polymeric micelles display brighter regions under UV-light illumination (yellow arrows in Figures 4c,d) corresponding to areas of higher accumulation of the luminescent probes. The intensity-to-temperature calibration curves of these regions recorded both in a suspension of DNPD-based polymeric micelles and in fixed MDA-MB-468 cells incubated with these micelles are analogous (Figure S19 in Supporting Information) demonstrating that the accumulation of the micelles does not impact the temperature dependence of the Eu³⁺ and Sm³⁺ emissions. Although the detailed analysis of the distribution of the nanoparticles within the cell lies out of the scope of the present work, and the identification of the spherical brighter areas would require the use of biological markers, from a morphological point of view these spots might correspond to the cells' nucleolus. To evaluate the effect of an external heat source in the temperature distribution at subcellular regions, we use a temperature controller attached to the imaging instrument described above. A total of 450 points were sampled (150 in the brighter and 300 in the darker regions) within the limits of 3 MDA-MB-468 cells for each temperature set between 296.45 and 303.55 K (Figure 5a-d). The histograms of both dark and bright regions display an inhomogeneous intracellular temperature progression up to ~20 degrees (Figure 5e-h and Figure S24 in Supporting Information), which is consistent with the previous reports on temperature mappings on living cells using diverse fluorophores as temperature probes 17, 33, 35, 36 (Table S8). Furthermore, all the histograms follow Gaussian distributions centered at values that increase with the external temperature set in the controller (Figure 5e-i). Remarkably, whereas for the darker regions the average temperatures are in excellent agreement with those set externally, for the brighter areas we observe an average temperature ~5 degrees higher than that of the rest of the cell (Figure 5i). The results in the suspensions of DNPD-based polymeric micelles and fixed MDA-MB-468 cells incubated with these micelles mentioned above dismiss the possibility of this temperature gradient be induced by the clustering of the micelles. Moreover, the observed temperature increment value is of the same order of magnitude than that already observed by Okabe *et al.*³⁰ and could be explained by the intense RNA transcription activity of the nucleolus. ^{46, 47}The real-time intracellular temperature imaging using these Ln³⁺-bearing polymeric micelles was recorded using the same procedure (Figure S21 in Supporting Information) to highlight the possibility of using the strategy described above for tracking thermal events occurring in subcellular level.

In summary, we reported here a unique system for real-time intracellular temperature mapping of living cells based on innovative luminescent Eu³⁺/Sm³⁺bearing polymeric micellar probes. The micelles and the Ln3+ complexes covalently bonded to its hydrophobic part were rationally designed to mitigate quenching effects induced by the culture medium. Moreover, the copolymerization strategy proposed permits easy incorporation of other functionalities, such as targeting different cell organelles, and the temperature is recorded from the ratio between the integrated areas of the ${}^4G_{5/2} \rightarrow {}^6H_{9/2}$ and ${}^5D_0 \rightarrow {}^7F_2$ transitions of the Sm³⁺ and Eu³⁺ ions, respectively, using a conventional (or minimally adapted) fluorescence microscope. A proof-of-concept experiment on breast metastatic adenocarcinoma cells (line MDA-MB-468) incubated with the Eu³⁺/Sm³⁺-bearing micelles under UV-light excitation yields 2D thermal images in the 296-304 K range with an uncertainty of 0.2 K. The thermal images allow us to delimitate the cells and to identify its nucleolus as brighter regions. By increasing the temperature of the culture medium, inhomogeneous intracellular temperature progressions up to ~20 degrees and an increment of ~5 degrees of the nucleolus temperature relative to the rest of the cell were discerned, which might be related to the activity of the thermo-generating organelles and highly exoenergetic



processes. The real-time intracellular temperature imaging using these Ln³+-bearing polymeric micelles opens new avenues to the detailed screening of thermal gradients within living cells due to endogenous or exogenous stimuli, providing important insights into the regulatory intracellular mechanisms, and thus contributing to an improved comprehension of the role of some subcellular thermo-generating organelles in the cellular functions.

Associated Content

Supporting Information: Materials and methods, Thermometric-imaging system, *In silico* experiments, Photoluminescence, Thermometric performance of the Ln³+-bearing micelles, Cell cultures, and Temperature imaging.

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Author Contributions

The manuscript was written through the contributions of all authors. RP synthetized the thermometric micelles, JZ recorded the photoluminescent measurements; CDSB wrote the MatLab routines, analyzed the microscopy images and the photoluminescent results, and draw the figures; YG performed the microscopy observations, ANCN and TES performed the in-silico experiments; PT and AM developed the cell temperature imaging experimental set-up; RM, PFS, AG, and LML performed the cell culture experiments; LDC and AM wrote the paper and co-supervised the project. All authors have approved the final version of the manuscript.

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Abbreviations

btfa - 4,4,4-trifluoro-1-phenyl-1,3-butanedione; DNPD - 1,3-di(naphthalen-2-yl)propane-1,3-dione; MPEGA - methoxy polyethylene glycol acrylate; PEGA - polyethylene glycol acrylate; MA - methyl acrylate; PhenA - 1,10-phenanthrolin-4-yl)methyl acrylate.

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Figures

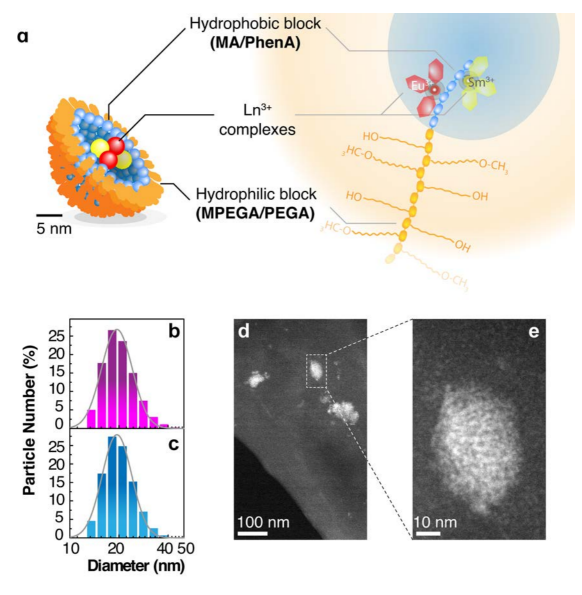


Figure 1. (a) Scheme of the Ln^{3+} -bearing polymeric micelles. The details of the hydrophobic and hydrophilic polymer chains are presented in Figure S1, while those of the Ln^{3+} complexes in Figures S4 and S5 in Supporting Information. Dynamic light scattering size distribution of the Ln^{3+} -bearing **(b)** btfa- and **(c)** DNPD- based polymeric micelles after filtration through a 100 nm membrane filter. The solid lines are the best fit to log-normal distributions ($r^2 > 0.99$) centered at 20.3±0.2 nm in both cases. **(d,e)** Scanning transmission electron microscope images of the Ln^{3+} -bearing btfa-based polymeric micelles.

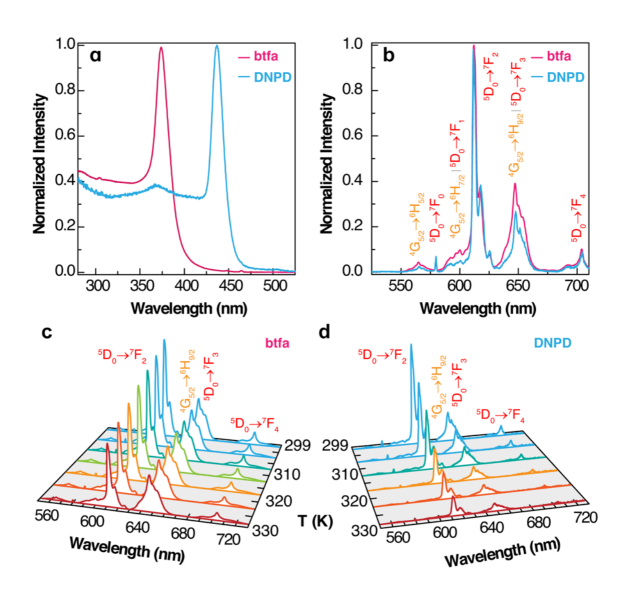


Figure 2. (a) Excitation spectra of btfa- and DNPD-based polymeric micelles at room temperature monitoring the ${}^5D_0 \rightarrow {}^7F_2$ transition of Eu³+. **(b)** Room temperature emission spectra of the btfa- and DNPD-based polymeric micelles. The transitions ascribed to Eu³+ and Sm³+ are signed in red and orange, respectively. Temperature dependence of the emission spectra recorded in the 299–328 K range for **(c)** btfa- and **(d)** DNPD-based polymeric micelles. In (b), (c) and (d) the excitation wavelengths used are 373 nm and 436 nm for btfa- and DNPD-based micelles, respectively.

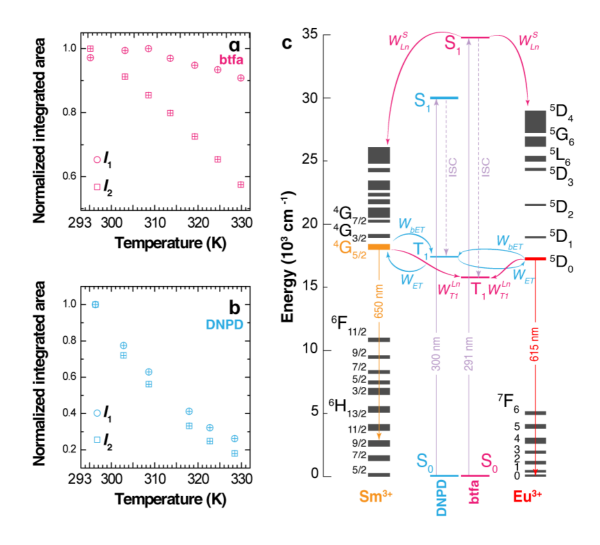


Figure 3. Temperature dependence of the integrated areas of the Sm³⁺ ${}^4G_{5/2} \rightarrow {}^6H_{9/2}$ (I_1) and the Eu³⁺ ${}^5D_0 \rightarrow {}^7F_2$ (I_2) transitions in the Ln³⁺-bearing (a) btfa- and (b) DNPD-based polymeric micelles. (c) Simplified energy level diagram for Sm³⁺, Eu³⁺, and ligands. The upwards arrow represents the initial absorptions by the ligands and ISC the intersystem crossing. For DNPD ligand, the forward intramolecular energy transfer rates (W_{ET}) are predominant via the $T_1 \rightarrow {}^4G_{5/2}$ (Sm³⁺) and $T_1 \rightarrow {}^5D_0$ (Eu³⁺) pathways. The backward energy transfer rates (W_{bET}) have a linear dependence with the temperature (Figure S6 in Supporting Information). For the btfa ligand, the forward intramolecular energy transfer rates (W_{Ln}^S) occur essentially *via* the S₁ $\rightarrow {}^4F_{5/2}$ (Sm³⁺) and S₁ $\rightarrow {}^5G_2$ (Eu³⁺) pathways. In this case, the luminescence quenching depends on the energy balance between the transfer processes (W^*), because the energy turns back to the ligand *via* the acceptor state T_1 .

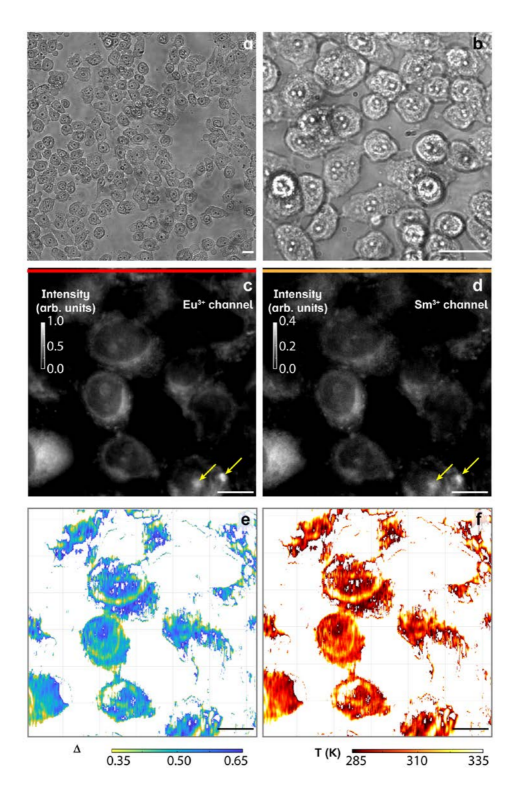


Figure 4. Room temperature microscopy images of MDA-MB-468 cells incubated with DNPD-based polymeric micelles under white-light illumination in (a) phase-contrast mode recorded with the color camera and (b) reflection mode recorded with the CMOS camera. Emission in the (c) Eu³⁺ and (d) Sm³⁺channels recorded with the CMOS camera and illuminating the cultures with a 365 nm LED. The yellow arrows sign illustrative brighter regions of the cells. Calculated 2D (e) Δ (using I_1 and I_2 channel images) and (f) temperature maps. All the scale bars correspond to 10 μm.

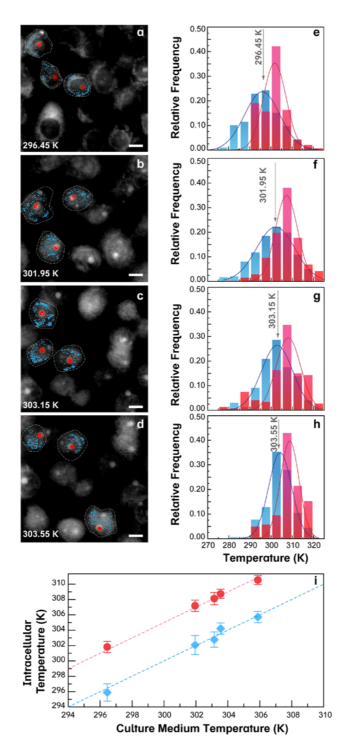


Figure 5. (a-d) Microscopy images (recorded in the Eu³⁺ channel) of the MDA-MB-468 cells incubated with DNPD-based polymeric micelles. The blue and red points mark, respectively, the locations used for temperature determination of the darker and brighter regions. The culture medium temperature values are indicated and The scale bars correspond to 10 μm. **(e-h)** Temperature histograms obtained from the selected points indicated in (a-d). The blue and red colors correspond to the blue (dark regions) and red (bright regions) points, respectively and the solid lines represent the best fits to Gaussian functions (r^2 >0.90). **(i)** Temperature determined using the intracellular thermometer (y) versus culture medium temperature (x) for the dark (blue) and bright (red) regions (e-h histograms). The blue line corresponds to y=x and the red one is shifted by 5 degrees (guides for the eyes). Data for 305.95 K is presented in Figure S16 in Supporting Information.

Real-time intracellular temperature imaging using lanthanide-bearing polymeric micelles

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Materials and methods

Materials

Selenium dioxide (SeO₂, 99.999%) was purchased from Strem Chemicals. Europium(III) chloride hexahydrate (EuCl₃(H₂O)₆, 99.99%), samarium(III) chloride hexahydrate (SmCl₃(H₂O)₆, 99.99%), 4,4,4-trifluoro-1-phenyl-1,3-butanedione (btfa, 99%) and 2-cyano-2-propyl dodecyl trithiocarbonate (97%, RAFT chain transfer agent) were all products of Sigma Aldrich and were used as received without further purification. 4-Methyl-1,10-phenanthroline was synthesized according to the method described by Belser *et al.*¹ The synthesis of 1,10-phenanthroline-4-carboxyaldehyde and 4-(hydroxymethyl)-1,10-phenantroline was accomplished following the straightforward methods reported in the literature with slight modifications.²⁻⁴

Methods

Synthesis of polymers

2,2'-Azobis(2-methylpropionitrile) (AIBN, Acros Organics, 98%) was recrystallized twice from ethanol. Methyl acrylate (MA, 99%, Aldrich) and poly(ethylene glycol) methyl ether acrylate (MPEGA) (Mn: 480 Da, Aldrich) was passed through a column of aluminum oxide to remove the inhibitors. Poly(ethylene glycol) acrylate (PEGA, Mn: 468 D, calculated by ¹H-NMR spectroscopy) was synthesized from polyethylene glycol 400 (Aldrich) and purified according to procedures previously described.⁵ Polymerizable ancillary ligand 4-(acryloyloxymethyl)-1,10-phenanthroline (PhenA) was prepared by esterification reaction of (3) with acryloyl chloride.

Macro chain transfer agent (CTA) P(MPEGA-co-PEGA) and the final diblock copolymer, P(PMEGA-co-PEGA)-b-P(MA-co-PhenA), were prepared using reversible addition-fragmentation chain transfer (RAFT) polymerization. In the preparation of the P(MPEGA-co-PEGA)-b-P(MA-co-PhenA), P(MPEGA-co-PEGA) was firstly synthesized using 2-cyano-2-propyl dodecyl trithiocarbonate as chain transfer agent, toluene as solvent, and AIBN as a radical source. The

polymerization of the final polymer was carried out in dioxane as solvent adapting the method used for the synthesis of the MacroCTA. The structure of the final polymer is shown in Figure S1.

Figure S1. Structure of the block copolymer used in the fabrication of the thermometric micelles.

Characterization of polymers

Nuclear Magnetic Resonance

Proton (1H), and carbon (13C) NMR and NMR spectra were recorded at room temperature in a BRUKER AV-400 spectrometer (400 MHz (1H), 100 MHz (13C) using CDCl3 or DMSO-d6 as solvent. 1H and 13C chemical shifts (δ) are reported in ppm and are referenced to the solvent peak the coupling constants (J) are given in Hz. 1H-NMR spectroscopy was used to confirm the synthesis of P(MPEGA-co-PEGA) and P(MPEGA-co-PEGA)-b-P(MA-PhenA) and determine the number average molecular weight, Mn (NMR), monomer percentage conversion, and composition of the polymers (degree of polymerization (DP), Mole fraction (X) and Weight fraction (Xw) for each monomer, **Table S1**). The calculated final molecular weight and the degree of polymerization were in concordance with the designed values.

Size exclusion chromatography

Molecular weights (Mn, Mw) and polydispersity (Mw/Mn) were measured by size exclusion chromatography using a Waters 2695 liquid chromatography system equipped with a Waters 2420 evaporation light scattering detector using a combination of two Styragel columns HR1 and HR4 calibrated using

poly(methylmethacrylate, PMMA) standards and THF as solvent at a flow rate of 1 mL per minute at 35 °C (308K).

Size-exclusion chromatograms (SEC) of both Macro Chain Transfer agent (CTA) P(MPEGA-co-PEGA) and the final diblock copolymer, P(PMEGA-co-PEGA)-b-P(MA-co-PhenA) showed a narrow polymer distribution indicating the formation of well-defined polymers with good polydispersity.

Molecular weight and molecular weight distribution and composition of the block copolymer and the macroCTA measured by ¹H NMR spectroscopy and SEC are listed in Table S1.

Table S1. Molecular weight and composition of the block copolymer P(MPEGA-co-PEGA)-b-P(MA-co-PhenA) and its macroCTA P(MPEGA-co-PEGA) determined by SEC or by NMR.

Polymer	Mn(Da) (NMR)	DP NMR	X (NMR)	Xw (NMR)	Mw(Da) (SEC)	Mn(Da) (SEC)	PDI (SEC)
P(MPEGA- co-PEGA)	13200	MPEGA (19)	MPEGA (0.70)	MPEGA (0.63)	9436 8752	8752	1.08
		PEGA (8) MPEGA (19)	PEGA (0.30) MPEGA (0.23)	PEGA (0.37) MPEGA (0.50)			
P(MPEGA- co-PEGA) -b- P(MA-co- PhenA)	18700	PEGA (8)	PEGA (0.11)	PEGA (0.20)	8803 8184	8184	1.07
		MA (50)	MA (0.61)	MA (0.23)			
		PhenA (4.5)	PhenA (0.05)	PhenA (0.06)			

Fourier-transform infrared spectroscopy

FT-IR spectra were recorded in a Perkin Elmer Spectrum 100 FT-IR spectrometer equipped with a universal attenuated total reflection sampling accessory. The spectra were collected over the 4000—380 cm⁻¹ range (attenuated total reflection correction performed).

The FTIR analysis confirmed the chemical structure of the block copolymer and the successful anchoring of the complex. The presence of lanthanide complexes was revealed by the presence of characteristic vibrational 1650-1500 cm⁻¹ bands associated with the metal-ligand coordination bonds. In the case of

the btfa complexes, the C-F stretching bands associated with the CF₃ groups of the ligand can also be observed at 1132 cm⁻¹. A comparison of the spectrum of polymer powders obtained by evaporation of the stock solutions in THF with that of micelles powders obtained by freeze-drying of the final micelle aqueous suspension showed no changes in the polymer bands, indicating that the integrity of complex was preserved during the auto self-assembly process of the polymer in water.

Differential scanning calorimetry

Thermal analysis was conducted by differential scanning calorimetry in a Q2000 TA equipment. The samples were quickly cooled down to −40°C (233 K) and then heated at the rate of 10 K·min⁻¹.

The DSC traces of the P(MPEGA-co-PEGA) MacroCTA were similar to those reported in the literature for P(MPEGA) homopolymers with a higher degree of polymerization. The crystallization of PEG side chains was observed at around - 39°C, and the melting peak was observed at nearly -12°C. However, the DSC traces of block copolymer showed neither crystallization peak nor Tg transition in the -40°C to 60°C temperature range indicating that both blocks are in a soft state at these temperatures.

Elemental analysis

Elemental analysis for C, H were performed using a Perkin Elmer CHN 2400 elemental analyzer with standard combustion conditions and handling of the samples in the air.

Synthesis of Ln3+-containing micelles

1,3-di(naphthalen-2-yl)propane-1,3-dione (DNPD) was prepared by adapting the synthetic method described by McGehee *et al.*⁶ Europium and samarium complexes, denoted by [Ln(L)₃(H₂O)₂] (Ln=Eu, Sm) were prepared following the synthetic procedure reported by Binnemans *et al.*⁷ The Ln(L)₃·2H₂O complexes were covalently attached by coordination with the ancillary ligand included in the polymer chain (PhenA) in the following way: the block copolymer and Ln(L)₃·2H₂O complexes were mixed in chloroform in stoichiometric proportion

(PhenA: Ln, 1:1; Eu(L)₃·2H₂O (7.5% mol) and Sm(L)₃·2H₂O (92.5% mol)) and heat overnight under reflux. After evaporation of the solvent, the polymer was dissolved in tetrahydrofuran (THF). Finally, the micelles were formed by self-assembly after the addition of water to the THF solution.

Polymeric micelles details

The hydrophilic block is a brush-like block of methoxy (MPEGA) and hydroxyl (PEGA) end-capped polyethylene glycol acrylates. The hydrophobic block is a methyl acrylate (MA) polymer containing a discrete number of the auxiliary ligand PhenA. The general formula of the lanthanide complexes is Ln(L1)₃L2 where L1 represents a β-diketonate ligand: 1,3-di(naphthalen-2-yl)propane-1,3-dione (DNPD) and L2 is an ancillary ligand, (1,10-phenanthrolin-4-yl)methyl acrylate (PhenA), a 1,10 phenanthroline derivative modified with a polymerizable acrylate group and pendant from the polymer chain after polymerization. The complexes are covalently attached to the polymer chain by coordination with the phenanthroline auxiliary ligand in chloroform. Then, the chloroform was evaporated, and the resulting product was dissolved in THF. Finally, the micelles are formed by self-assembly after the addition of water to the THF solution. The hydrophilic PEG block forms the outer shell of the micelles and the non-watersoluble methyl acrylate block containing the lanthanide complexes forms the inner shell. The presence of the PEG block provides simultaneously stability in aqueous media, biocompatibility, enhanced cellular uptake, and low toxicity to the nanostructures. Several batches of the btfa- and DNPD-based micelles were prepared with analogous structural and spectroscopic properties.

Chemical-physical characterization of the micellar solutions

Inductively coupled plasma optical emission spectrometry

Analysis of thermometric micelles by inductively coupled plasma optical emission spectrometry in a plasma 40 ICP Perkin-Elmer spectrometer yielded a concentration of Eu and Sm in the micelle aqueous dispersion of 0.44×10⁻⁴ mol/L, and 4.60×10⁻⁴ mol/L, respectively, corresponding to an Eu/Sm molar ratio of 0.095. The samples for analysis were prepared by digestion of 1 mL of the micelle

aqueous dispersion with concentrated HNO₃ (1 mL) overnight and then diluting to a final volume of 10 mL with ultrapure water.

Dynamic light scattering and zeta potential

Dynamic light scattering (DLS) and zeta potential measurements of the ferrofluids were performed on a Malvern Zetasizer NS (Malvern Instruments Ltd., Worcestershire, UK) using a He–Ne laser with a 633 nm wavelength and a detector angle of 173°. The measurements were repeated three times, and the average was calculated.

<u>Transmission electron microscopy</u>

Transmission electron microscopy (TEM) imaging was performed in a FEI Tecnai F30 microscope equipped with an EDAX XEDS system with an energy resolution of 130 eV. Samples for TEM analysis were prepared by dip-coating on lacey carbon grids.

Energy dispersive X-ray spectroscopy (EDS) of Sm and Eu content was performed in selected areas within the thermometric micelles. The presence of $SmM_{\alpha 1}$ $SmL_{\alpha 1}$ and $SmL_{\beta 1,2}$ peaks was observed (Figure S2).

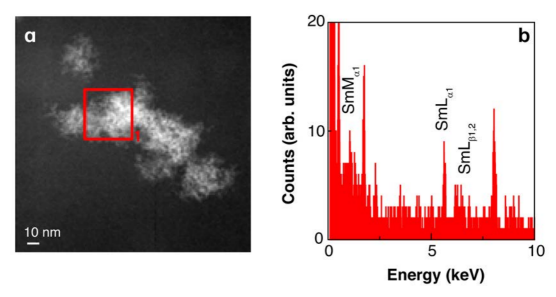


Figure S2. EDS analysis on an area of a sample prepared by evaporation of the ferrofluid on a carbon-coated copper grid. The Sm emission can be clearly distinguished.

Temperature-dependent photoluminescence

The photoluminescence spectra were measured in the room-temperature 300 –328 K range with a modular double-grating excitation spectrofluorimeter with a TRIAX 320 emission monochromator (Fluorolog-3, Horiba Scientific) coupled to an R928 Hamamatsu photomultiplier using the front face acquisition mode. The excitation source was a 450 W Xe arc lamp. For the temperature-dependent experiments, the samples were in contact with a heating stage (Linkam, THMS 600).

Stability of the micellar thermometric probes

To ensure a firm attachment of the lanthanide emitters to the copolymer forming the micelles, we followed the strategy of fixing the lanthanide Ln(L1)₃ complexes to the polymer backbone by covalent binding the complex (instead of physical absorption) with a strong lanthanide ligand, as phenanthroline, that is incorporated in the polymer structure. This strategy also produces an enhancement of the emission intensity, with respect to the pure Ln(L1)₃ complexes.

The attachment of the Ln(L)₃ complexes to the phenanthroline ancillary ligand in the polymer was studied by Fourier transform infrared spectroscopy (FTIR) (Figure S3). The disappearance of the broad absorption bands centered at 3380 cm⁻¹, corresponding to the H₂O coordination molecules in the Ln(L)₃(H₂O)₂ complexes, indicates the displacement of these molecules by coordination of the lanthanides to the phenanthroline ligand in both types of micelles. In the case of the btfa micelles, the FTIR spectrum shows a relative increase of the intensity of the (C=O) stretching band at 1610 cm⁻¹, with respect to that at 1597cm⁻¹, and a widening of the vibration frequency band at 1530 cm⁻¹ typical of pure Ln(btfa)₃Phen complexes. In the case of DNPD micelles, an increase in the strength of the band at 1495 cm⁻¹, and a shift of the 1375 cm⁻¹ band to higher wavenumbers (1404 cm⁻¹) are also evidence of lanthanide phenanthroline coordination bonds.

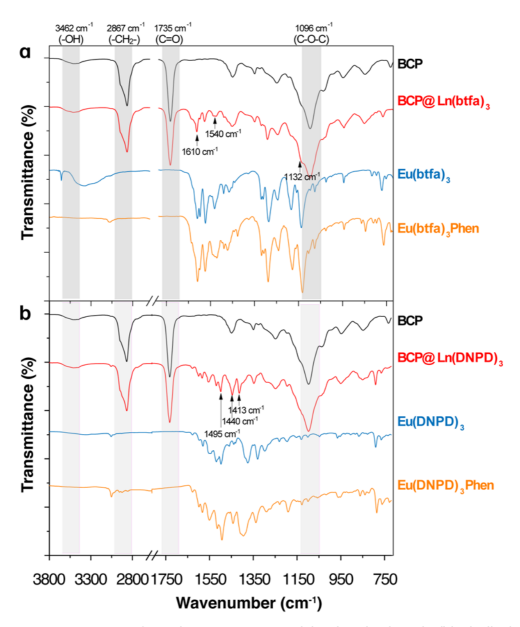


Figure S3. FTIR spectra for P(MPEGA-co-PEGA)-b-P(MA-b-PhenA) (black line), P(MPEGA-co-PEGA)-b-P(MA-b-(PhenA@EuSm(L)₃) (red line), Eu(L)₃(H₂O)₂ complex (blue line) and Eu(L)₃Phen (orange line) for **(a)** L=btfa and **(b)** L=DNPD based material.

Lanthanide metal content in the micellar suspension was determined by Inductively Coupled Plasma Atomic Emission Spectrometry (ICP-AES) in a plasma 40 ICP Perkin-Elmer spectrometer. Samples for ICP-AES were prepared by digestion of 1 mL of the micellar solution after freeze-drying with concentrated HNO₃ overnight and diluted with ultrapure water. The experimental ratios of the Ln³⁺ ions were 2.42% and 2.84% w/w, for DNDP and btfa complexes, respectively. Considering the nominal values of 2.84% and 3.03% w/w used in the encapsulation reaction, the loading efficiency was 85% and 94% for DNDP

and btfa complexes, respectively. The stability of the micelles suspensions was studied by DLS (Figure S4). It was observed that the size remains unchanged at least 6 months after storage.

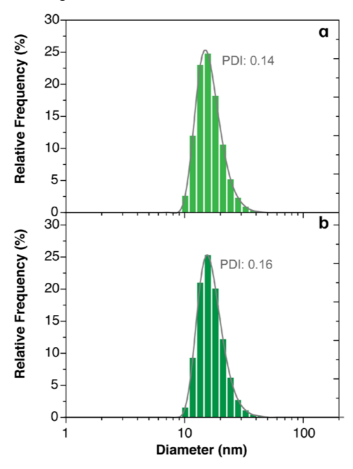


Figure S4 Histograms of the hydrodynamic diameter for btfa-based micelles measured **(a)** after 1 day and **(b)** after 6 months of the preparation.

The effect of dilution on the structural and optical stabilities of the micellar solutions was attested by DLS (Figure S5a) and photoluminescence measurements (Figure S5b). Dilutions in ultrapure water of the as-prepared solution by 1/5 and 1/10 did not cause any change neither in the size distribution nor in the emission spectra of the micelles. Moreover, the relative ratio of Eu³⁺/Sm³⁺ intensities was also maintained after dilution.

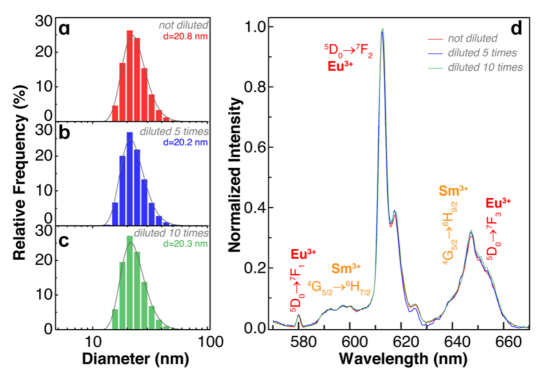


Figure S5 (a) Histograms of the hydrodynamic diameter (d) and **(b)** room-temperature emission spectra of btfa-based micelles under 365 nm excitation for different dilution factors.

The leakage of the btfa- and DNPD-based complexes from the micelles was checked using cryo-TEM. The fast freezing of the suspension in this technique ensures that the frozen sample precisely reproducing the state of the materials in the liquid. Cryo-STEM images together with electron energy loss spectroscopy (EELS) and EDS analysis of micelle samples several weeks after preparation showed no presence of lanthanides outside the micelles, demonstrating the absence of significant leaking from the interior of the micelles into the bulk of the suspension.

The optical properties of the Ln³+-bearing complexes changed in cell culture media with respect to pure water. It has experimentally observed that Ln(L1)₃-2H₂O complexes lost their emission in the cell culture medium. The linking the Ln(L1)₃-2H₂O complexes to the phenanthroline ligand protects from luminescence quenching effects by steric shielding. However, it was observed that the shielding effect was not sufficient in the case of Ln(btfa)₃Phen that decreased gradually its emission intensity when transferred from water to the cell culture medium. Nevertheless, when the bfta ligands were replaced by DNPD the luminescence was preserved in cell culture medium and persisted after

internalization of the micelles in the cells. Such enhancement of the emission stability in the cell culture medium can be attributed to the improvement of the steric effects of the coordination shell due to the larger volume of the DNPD ligand as compared to the btfa one (Figure S6).

$$\begin{array}{c|c} \mathbf{G} & \mathbf{b} \\ \mathbf{H_2O} & \mathbf{O} \\ \mathbf{H_2O} & \mathbf{O} \\ \mathbf{H_2O} & \mathbf{O} \\ \mathbf{H_2O} & \mathbf{O} \\ \mathbf{CF_3} \\ \mathbf{In} \\ \mathbf{Ln} \\ \mathbf{DNPD})_3 (\mathbf{H_2O})_2 \end{array}$$

Figure S6. Structure of **(a)** Ln(DNPD)₃(H₂O)₂ and **(b)** Ln(btfa)₃(H₂O)₂ complexes.

Thermometric-imaging system

The fluorescence microscope used in cell imaging experiments was a Leica DMI3000B equipped with a DFC 7000 color camera. The pixel-to-pixel lateral resolution is ~97 nm, although the ultimate spatial resolution is determined by the microscope optics, about 1 μ m. The beam splitter and the CMOS camera are from Hamamatsu instrument and a Leica fluorescence microscope incorporating a Gemini beam splitter and a CMOS Orca 4.0 camera (both from Hamamatsu) connected to a computer endowed with a LabView image-processing program. The beam splitter was equipped with a Semrock FF635-Di01 dichroic, an FBH650-40 bandpass filter, central wavelength 650 nm, full-width at half maximum, fwhm=40 nm (for Sm³+ emission), and a Chroma ET610/20 NC491994 bandpass filter, central wavelength 610 nm, fwhm=20 nm (for Eu³+ emission).

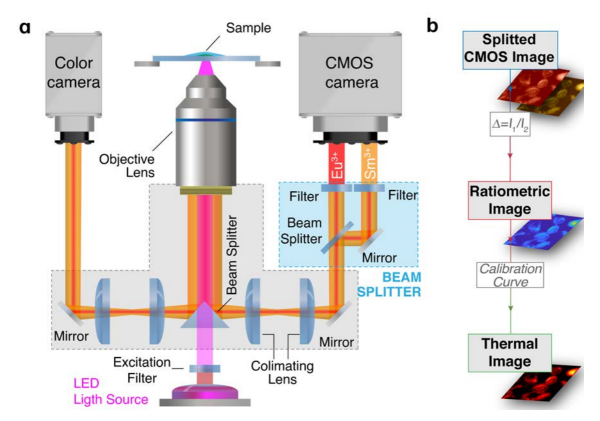


Figure S7. (a) Scheme of the microscope setup used to acquire simultaneously the regular color image (DFC 7000 color camera) and the split CMOS image (Orca 4.0 camera) for temperature mapping. The LED light (365 nm) passes the excitation filter before illuminating the sample. The dichroic mirror blocks the excitation light to prevent CMOS damage The emitted light is divided by a customized beam-splitter system that projects the Sm³+ (620–660 nm) and the Eu³+ (600–620 nm) spectral regions in each half of the CMOS camera detector. **(b)** Flux diagram of the image processing using the MatLab® routine.

In silico experiments

Since there are no Gd^{3+} (or even La^{3+} or Lu^{3+}) samples to obtain the ligands excited energies T_1 and S_1 , quantum chemistry calculations are useful to estimate these states. In this work, DFT and TD-DFT (time-dependent DFT) calculations, using Gaussian 09 program,⁸ were performed in the presence of a solvent (water) within the Polarizable Continuum Model (PCM) using the integral equation formalism variant (IEFPCM).⁹⁻¹¹ For both, structure optimizations and excited states calculations were used ω B97-XD functional.¹² Due to a large number of atoms (> 200), the basis set STO-3G (for C, H, N and O atoms) was used while the Ln^{3+} ions were treated with MWB52 (Eu^{3+}) and MWB51 (Sm^{3+}).¹³

There is no significant difference in structures between Eu³+–DNPD and Sm³+–DNPD coordination compounds. Figure S8a shows the structure from DFT simulations, the highlighted atoms (white spheres) represent the ancillary ligand bonded with the polymer. Figure S8b-c shows the main compositions of molecular orbitals (from occupied to unoccupied ones) involved in the T¹ excited state.

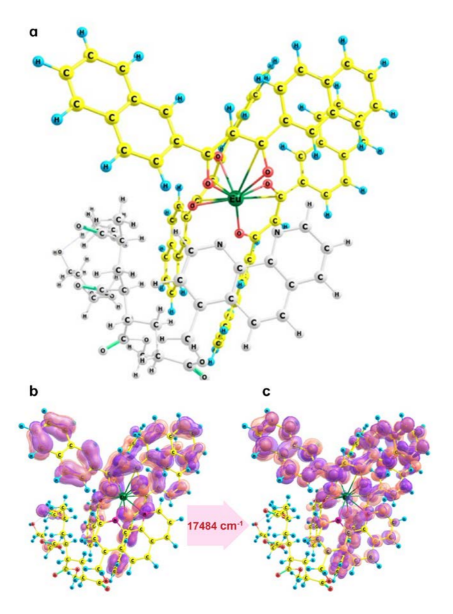


Figure S8. (a) Optimized structure for the DNPD-Ln³+ complexes obtained by DFT calculations (ω B97-XD/MWB52(Eu³+)/MWB51(Sm³+)/STO-3G level of theory). The colorless atoms represent the ancillary ligand (1,10-Phenanthroline + a small polymeric part). (b, c) TD-DFT calculations (in the same level of theory) showing the main molecular orbitals contributions (isosurfaces of 0.03 e/a_0^3) for the T¹ state. It is noticed that there is no significant participation of the ancillary ligand part in the T¹ state, this state is determined predominantly by DNPD ligands.

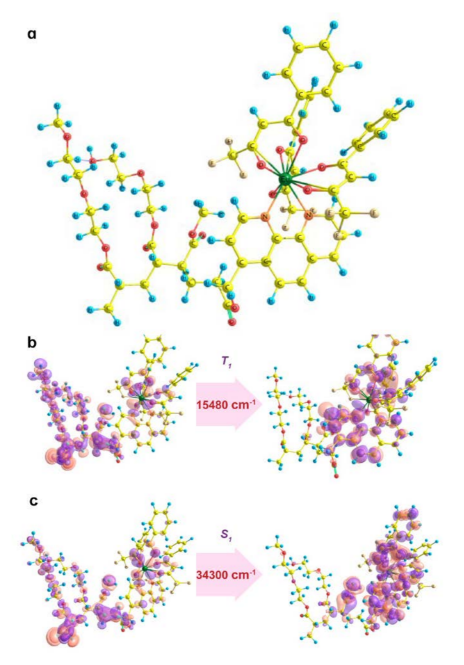


Figure S9. (a) Optimized structure for the btfa-Ln³+ complexes obtained by DFT calculations (ω B97-XD/MWB52(Eu³+)/MWB51(Sm³+)/STO-3G level of theory). (b, c) TD-DFT calculations (in the same level of theory) showing the main molecular orbitals contributions (isosurfaces of 0.03 e/a_0^3) for the T¹ and S¹ states. Both states are formed predominantly by an electronic density flux from the occupied orbitals close to the polymer part to unoccupied orbitals close to the btfa ligands.

Intramolecular energy transfer

The intramolecular energy transfer (IET) rates were calculated taking into account the dipole-dipole (W_{d-d}), dipole-multipole (W_{d-m}), and exchange mechanisms (W_{ex}) in coordination compounds, ¹⁴⁻¹⁷ Eqs.S1-S3. The coordination compounds herein analyzed are based on structural data and the excited states obtained from TD-DFT calculations. More details on the quantities in the equations below are in Refs. 14-19.

The IET rates can be obtained by using multipolar and exchange interactions. Eq. S1 stands for the dipole-multipole (4-, 16- and 64-poles) contributions, K = 2, 4, and 6, respectively.

$$W_{d-m} = \frac{S_L}{(2J+1)G} \frac{2\pi e^2}{\hbar} \sum_K (K+1) \frac{\langle r^K \rangle^2}{(R_L^{K+2})^2} \langle f \| C^{(K)} \| f \rangle^2 (1 - \sigma_K)^2 \times \langle \psi^* J^* \| U^{(K)} \| \psi J \rangle^2 F$$
(S 1)

where S_L is the dipole strength of the ligand transition concerning in IET, $\langle r^K \rangle$ are the 4f radial integrals, G is the ligand state degeneracy, R_L (equals to 6.6 and 5.9 Å for the DNPD and btfa-based complexes, respectively) is the donor-acceptor states distance, $\langle f || C^{(K)} || f \rangle$ is the reduced matrix element of Racah's tensor operator, $(1 - \sigma_K)$ are the shielding factor, and $\langle \psi^* J^* || U^{(K)} || \psi J \rangle^2$ are the squared reduced matrix elements and their values are tabulated in Ref. 20.

For the dipole-dipole contribution, the Eq. S2 is used:

$$W_{d-d} = \frac{S_L (1 - \sigma_1)^2}{(2J + 1)G} \frac{4\pi}{\hbar} \frac{e^2}{R_L^6} \sum_{\lambda} \Omega_K^{FED} \langle \psi^* J^* \| U^{(K)} \| \psi J \rangle^2 F$$
 (S 2)

where the intensity parameters Ω_K^{FED} are obtained by using only the forced electric dipole mechanism (original Judd-Ofelt theory^{21, 22}) with the Simple Overlap model.^{23, 24}

The energy transfer rate by the exchange interaction (Eq. S3) can be calculated by:

$$W_{ex} = \frac{\langle 4f | L \rangle^4}{(2J+1)} \frac{8\pi}{3\hbar} \frac{e^2}{R_L^4} \langle \psi^* J^* || S || \psi J \rangle^2 \sum_m \left| \langle \phi | \sum_j \mu_z(j) s_m(j) | \phi^* \rangle \right|^2 F$$
 (S 3)

where $\langle 4f|L\rangle$ is the overlap integral between the ligand donor state and the 4f acceptor state at R_L distance, s_m the spin operator of electron j in the ligand, μ_Z is the dipole operator (z-component), and $\langle \psi^*J^*||S||\psi J\rangle$ is the reduced matrix elements of the spin operator, which were obtained using free-ion wavefunctions in the intermediate coupling scheme. ²⁵

The spectral overlap F is related to the energy mismatch condition and contains a sum over Franck-Condon factors. ¹⁵ In the case of energy transfer involving lanthanide complex, F can be obtained by:

$$F = \frac{1}{\hbar \gamma_L} \sqrt{\frac{\ln(2)}{\pi}} e^{-\left(\frac{\Delta}{\hbar \gamma_L}\right)^2 \ln(2)}$$
 (S 4)

The Eq. S4 can be assumed only when the corresponding bandwidth at half-height for the ligand (γ_L) is much larger than the lanthanide (γ_{Ln}) ion, $\gamma_L \gg \gamma_{Ln}$. The Δ (in cm⁻¹) is the band maximum energy difference between ligand donor state and lanthanide ion acceptor state, $\Delta E = E_L - E_{Ln}$. The forward energy transfer rates (W_{ET}) can be calculated by the sum over all Eq. S1, Eq. S2 and Eq. S3 in the same pathway:

$$W_{ET} = W_{d-m} + W_{d-d} + W_{ex} {(S 5)}$$

The backward energy transfer rates (W_{bET}), that is, the energy returned from acceptor to donor state, is obtained with the same above equations, except for multiplying the spectral overlap F (Eq. S4) by the Boltzmann's factor:

$$F_{bET} = \frac{1}{\hbar \gamma_L} \sqrt{\frac{\ln(2)}{\pi}} e^{-\left(\frac{\Delta E}{\hbar \gamma_L}\right)^2 \ln(2)} \times e^{-\left(\frac{|\Delta E|}{k_B T}\right)}$$
 (S 6)

where k_B is the Boltzmann's constant.

Energy transfer pathways

The pathways and its predominant mechanism to the forward intramolecular energy transfer rates (W_{ET}) for the complexes inside the polymeric micelle structure are shown in Table S2 and Table S3 for DNPD-based polymeric micelles and Table S5 and Table S6 for btfa-based micelles. It was analyzed different energy transfer pathways in which ligand-to-metal energy transfer can occur, from both T₁ (17500 cm⁻¹ and 15480 cm⁻¹) or S₁ (30000 cm⁻¹ and 34300 cm⁻¹) ligand states for DNPD and BTFA, respectively.

Table S2. Intramolecular energy transfer rates (in s⁻¹) for Eu³⁺–DNPD complex in polymeric micelles. The donor-acceptor distance is R_L =6.6 Å.

Pathway	Donor State	Acceptor State	W _{ET}	Mechanism
1	T_1	$^{7}F_{1} \rightarrow {}^{5}D_{0}$	1.972×10 ⁶	exchange
2	T_1	$^{7}\text{F}_{0} \rightarrow {}^{5}\text{D}_{0}$	0.447	dipole-multipole*
3	T_1	$^{7}\text{F}_{1} \rightarrow {}^{5}\text{D}_{1}$	1.734	exchange
4	T_1	$^7F_0 \rightarrow {}^5D_1$	1.504×10^3	exchange
5	T_1	$^7F_1 \rightarrow {}^5D_2$	1.236×10 ⁻³	exchange
6	T_1	$^{7}\text{F}_{0} \rightarrow {}^{5}\text{D}_{2}$	4.191×10 ⁻⁷	dipole-multipole
7	S_1	$^{7}F_{1} \rightarrow {}^{5}D_{0}$	1.898×10^3	exchange
8	S_1	$^{7}\text{F}_{0} \rightarrow {}^{5}\text{D}_{0}$	12.39	dipole-multipole*
9	S_1	$^{7}F_{1} \rightarrow {}^{5}D_{1}$	2.857×10^3	dipole-multipole
10	S_1	$^{7}F_{0} \rightarrow {}^{5}D_{1}$	2.319×10 ⁴	exchange
11	S_1	$^7F_1 \rightarrow {}^5D_2$	4.109×10^3	exchange
12	S_1	$^{7}\text{F}_{0}$ \rightarrow $^{5}\text{D}_{2}$	2.313×10 ⁵	dipole-multipole
13	S_1	$^{7}\text{F}_{0} \rightarrow {}^{5}\text{D}_{3}$	0.000	none
14	S_1	$^{7}\text{F}_{1} \rightarrow {}^{5}\text{D}_{3}$	1.108×10 ⁵	dipole-multipole
15	S_1	$^7F_0 \rightarrow ^5L_6$	3.170×10 ⁴	dipole-dipole
16	S_1	$^7F_1 \rightarrow {}^5L_6$	2.773×10^3	dipole-dipole
17	S_1	$^7F_0 \rightarrow ^5D_4$	1.368×10^3	dipole-dipole
		Total	2.383×10 ⁶	

^{*} Results obtained considering the J-mixing effect.

Table S3. Intramolecular energy transfer rates (in s⁻¹) for Sm³⁺–DNPD complex in polymeric micelles. The donor-acceptor distance is R_L =6.6 Å.

Pathway	Donor State	Acceptor State	W _{ET}	Mechanism
1	T_1	$^6 H_{5/2} ightarrow {}^4 G_{5/2}$	3.053×10 ⁵	exchange
2	T_1	$^6H_{5/2} \rightarrow {}^4F_{3/2}$	0.000	none
3	T_1	$^6H_{5/2} \rightarrow {}^4G_{7/2}$	5.059×10 ⁻²	exchange
4	T_1	$^{6}\text{H}_{5/2} ightarrow {}^{4}\text{I}_{9/2}$	9.471×10 ⁻⁶	dipole-multipole
5	T_1	$^6 H_{5/2} ightarrow {}^4 M_{15/2}$	2.641×10 ⁻⁷	dipole-dipole
6	T_1	$^{6}\text{H}_{5/2} \rightarrow {}^{4}\text{I}_{11/2}$	1.785×10 ⁻⁸	dipole-dipole
7	T_1	$^{6}\text{H}_{5/2} ightarrow {}^{4}\text{I}_{13/2}$	3.286×10 ⁻⁹	dipole-dipole
8	T_1	$^{6}\text{H}_{5/2} ightarrow {}^{4}\text{F}_{5/2}$	1.246×10 ⁻⁶	exchange
9	T_1	$^6 H_{5/2} \rightarrow {}^4 M_{17/2}$	4.463×10 ⁻¹²	dipole-dipole
10	T_1	$^6H_{5/2} \to {}^4G_{9/2}$	7.126×10 ⁻¹³	dipole-dipole
11	S_1	$^6H_{5/2} ightarrow {}^4G_{5/2}$	0.687	exchange
12	S_1	$^{6}H_{5/2} \rightarrow {}^{4}F_{3/2}$	0.000	None
13	S_1	$^6H_{5/2} \rightarrow {}^4G_{7/2}$	1.308×10^3	dipole-multipole
14	S_1	$^{6}\text{H}_{5/2} \rightarrow {}^{4}\text{I}_{9/2}$	1.837×10 ⁴	dipole-multipole
15	S_1	$^{6}H_{5/2} \rightarrow {}^{4}M_{15/2}$	1.530×10^3	dipole-dipole
16	S_1	$^{6}\text{H}_{5/2} \rightarrow {}^{4}\text{I}_{11/2}$	5.114×10 ²	dipole-dipole
17	S_1	$^{6}H_{5/2} \rightarrow {}^{4}I_{13/2}$	3.668×10^3	dipole-dipole
18	S_1	$^6 H_{5/2} \rightarrow {}^4 F_{5/2}$	3.747×10^2	exchange
19	S_1	$^6H_{5/2} ightarrow {}^4M_{17/2}$	2.643×10^3	dipole-dipole
20	S_1	$^6H_{5/2} \rightarrow {}^4G_{9/2}$	2.332×10^3	dipole-dipole
		Total	3.36×10⁵	

Pathway 1, for both Eu³⁺ and Sm³⁺ as acceptors, is the dominant one and corresponds to ~83% and ~91% of the total forward IET rates, respectively. The backward energies transfer rates (W_{bET}) from pathways 1 are the keys to explain why the intensities decrease when the temperature grows. Indeed, Figure S10. shows the increase in W_{bET} when the temperature increases, depopulating the emitting levels 5D_0 (Eu³⁺) and $^4G_{5/2}$ (Sm³⁺). W_{bET} for Eu–DNPD is more affected by the temperature than the Sm–DNPD one, explaining why the integrated area (Figure 3b) of $^5D_0 \rightarrow ^7F_2$ transition decay faster than the integrated area of $^4G_{5/2} \rightarrow ^6H_{9/2}$ one, as pointed in Figure S10..

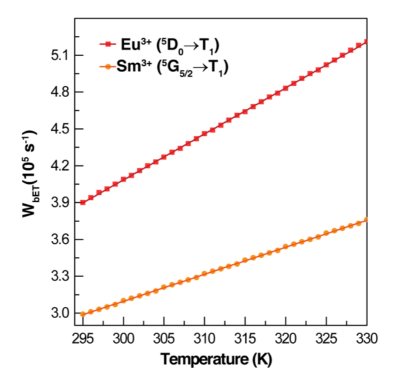


Figure S10. Backward IET rates from the emitters levels of Eu^{3+} (5D_0) and Sm^{3+} ($^4G_{5/2}$) ions for DNPD-based lanthanide complexes. The rate $^5D_0 \to T_1$ grows faster than the $^4G_{5/2} \to T_1$, resulting in a more effective depopulation on 5D_0 level, therefore, the integrated area of the $^5D_0 \to ^7F_2$ transition decreases faster than the integrated area of the $^5G_{5/2} \to ^6H_{9/2}$. The dependences were fitted by straight lines (Table S4).

Table S4. Linear coefficients and their respective uncertainties for the backward IET rates tendencies of Figure S10.

lon	Intercept (s ⁻¹)	Slope (10 ⁻² s ⁻¹ ·K ⁻¹)	r²
Eu ³⁺	-7.15±0.02	3.746±0.007	0.99
Sm ³⁺	-3.50 ± 0.02	2.198±0.005	0.99

Table S5. Intramolecular energy transfer rates (in s^{-1}) for Eu³⁺–btfa complex in polymeric micelles. The donor-acceptor distance is R_L =5.9 Å.

Pathway	Donor State	Acceptor State	W _{ET}	Mechanism
1	T_1	$^{7}F_{1} \rightarrow {}^{5}D_{0}$	6.453×10 ³	exchange
2	T_1	$^{7}F_{0}$ $ ightarrow$ $^{5}D_{0}$	1.671×10 ⁻⁴	dipole-multipole*
3	T_1	$^{7}\text{F}_{1} \rightarrow {}^{5}\text{D}_{1}$	2.788×10 ⁻⁴	exchange
4	T_1	$^{7}\text{F}_{0} \rightarrow {}^{5}\text{D}_{1}$	0.229	exchange
5	T_1	$^{7}\text{F}_{1} \rightarrow {}^{5}\text{D}_{2}$	1.316×10 ⁻⁷	exchange
6	T_1	$^7F_0 o {}^5D_2$	3.035×10 ⁻¹¹	dipole-multipole
7	\mathcal{S}_1	$^{7}\text{F}_{1} \rightarrow {}^{5}\text{D}_{0}$	22.77	exchange
8	S_1	$^{7}\text{F}_{0} \rightarrow {}^{5}\text{D}_{0}$	0.108	dipole-multipole
9	S_1	$^{7}\text{F}_{1} \rightarrow {}^{5}\text{D}_{1}$	42.48	dipole-multipole
10	S_1	$^{7}F_{0} \rightarrow {}^{5}D_{1}$	6.084×10^2	exchange
11	\mathcal{S}_1	$^{7}\text{F}_{1} \rightarrow {}^{5}\text{D}_{2}$	2.339×10^{2}	exchange
12	S_1	$^{7}\text{F}_{0}$ \rightarrow $^{5}\text{D}_{2}$	1.014×10 ⁴	dipole-multipole
13	S_1	$^{7}\text{F}_{0} \rightarrow {}^{5}\text{D}_{3}$	0.000	none
14	S_1	$^{7}\text{F}_{1} \rightarrow {}^{5}\text{D}_{3}$	2.739×10^3	dipole-multipole
15	\mathcal{S}_1	$^{7}\text{F}_{0} \rightarrow {}^{5}\text{L}_{6}$	4.839×10^3	dipole-dipole
16	S_1	$^{7}F_{1} \rightarrow {}^{5}L_{6}$	3.680×10^{2}	dipole-dipole
17	S_1	$^{7}F_{0}$ $ ightarrow$ $^{5}D_{4}$	1.714×10^3	dipole-dipole
18	\mathcal{S}_1	$^7F_1 \rightarrow {}^5G_2$	6.128×10 ⁴	exchange
19	S_1	$^{7}F_{1} \rightarrow {}^{5}G_{3}$	8.721×10 ³	dipole-multipole
		Total	9.716×10 ⁴	

^{*} Results obtained considering the J-mixing effect.

Table S6. Intramolecular energy transfer rates (in s^{-1}) for Sm³⁺-btfa complex in polymeric micelles. The donor-acceptor distance is R_L =5.9 Å.

Pathway	Donor State	Acceptor State	W _{ET}	Mechanism
1	T_1	$^6 H_{5/2} \rightarrow {}^4 G_{5/2}$	2.559×10 ⁵	exchange
2	T_1	$^{6}H_{5/2} \rightarrow {}^{4}F_{3/2}$	0.000	none
3	T_1	$^6H_{5/2} ightarrow {}^4G_{7/2}$	6.456×10 ⁻²	exchange
4	T_1	$^{6}H_{5/2} \rightarrow {}^{4}I_{9/2}$	8.037×10 ⁻¹⁰	dipole-multipole
5	T_1	$^6H_{5/2} ightarrow {}^4M_{15/2}$	1.726×10 ⁻¹¹	dipole-dipole
6	T_1	$^{6}H_{5/2} \rightarrow {}^{4}I_{11/2}$	1.104×10 ⁻¹²	dipole-dipole
7	T_1	$^{6}\text{H}_{5/2} ightarrow {}^{4}\text{I}_{13/2}$	1.904×10 ⁻¹³	dipole-dipole
8	T_{1}	$^{6}\text{H}_{5/2} \rightarrow {}^{4}\text{F}_{5/2}$	1.097×10 ⁻⁶	exchange
9	T_1	$^{6}H_{5/2} \rightarrow {}^{4}M_{17/2}$	2.140×10 ⁻¹⁶	dipole-dipole
10	T_1	$^6H_{5/2} \to {}^4G_{9/2}$	3.420×10 ⁻¹⁷	dipole-dipole
11	\mathcal{S}_1	$^6H_{5/2} \rightarrow {}^4G_{5/2}$	1.999×10 ⁴	exchange
12	\mathcal{S}_1	$^{6}\text{H}_{5/2} ightarrow {}^{4}\text{F}_{3/2}$	0.000	none
13	\mathcal{S}_1	$^6H_{5/2} \rightarrow {}^4G_{7/2}$	2.350×10 ⁵	exchange
14	\mathcal{S}_1	$^{6}\text{H}_{5/2} ightarrow {}^{4}\text{I}_{9/2}$	6.433×10^2	dipole-multipole
15	\mathcal{S}_1	$^6 H_{5/2} ightarrow {}^4 M_{15/2}$	46.82	dipole-dipole
16	\mathcal{S}_1	$^{6}\text{H}_{5/2} ightarrow {}^{4}\text{I}_{11/2}$	14.70	dipole-dipole
17	\mathcal{S}_1	$^{6}\text{H}_{5/2} ightarrow {}^{4}\text{I}_{13/2}$	97.03	dipole-dipole
18	\mathcal{S}_1	$^{6}\text{H}_{5/2} ightarrow {}^{4}\text{F}_{5/2}$	3.637×10^6	exchange
19	\mathcal{S}_1	$^6 H_{5/2} ightarrow {}^4 M_{17/2}$	57.44	dipole-dipole
20	\mathcal{S}_{1}	$^6H_{5/2} o {}^4G_{9/2}$	4.536×10^3	dipole-dipole
		Total	4.153×10 ⁶	

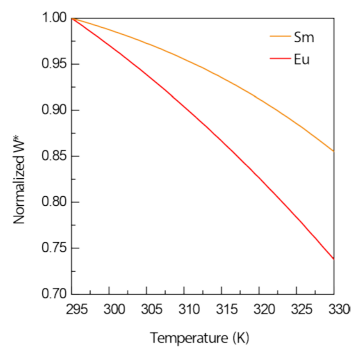


Figure S11. Temperature dependence of the normalized W* for btfa-based lanthanide complexes. The energy transfer from the Eu³+ emitting level to the T_1 state is in the same order of magnitude of the energy transfer from S_1 to the Eu³+ higher states. In the case of Sm³+, the $W_{T_1}^{Ln}$ is one order of magnitude lower than the W_{Ln}^{S} . This corroborates with the temperature dependence of the integrated areas as shown in Figure 3a.

Photoluminescence

Room temperature excitation

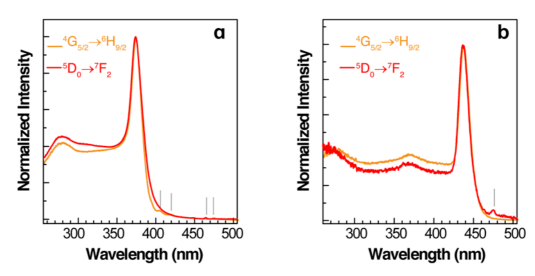


Figure S12. Room-temperature excitation spectra monitoring ${}^5D_0 \rightarrow {}^7F_2$ transition of Eu³⁺ and ${}^4G_{5/2} \rightarrow {}^6H_{9/2}$ transition of Sm³⁺ in **(a)** btfa and **(b)** DNPD-based micelles. The corresponding absorption spectra are presented in **(c)** and **(d)**, respectively.

Regardless of the monitored emission line of Eu³⁺ (or Sm³⁺) in both polymeric micelles at broadband is observed in excitation spectra with a maximum around 380 nm for btfa-based polymeric micelles and 440 nm for DNPD-based material. These absorption bands are ascribed to the ligand absorption.

The emission spectra in the physiological temperature range were recorded for thermometric analysis. Firstly, a straight baseline was removed for each spectrum and the conversion from wavelength to energy scale was done. Also the intensity as a function of wavelength, $f(\lambda)$, in the wavelength range $d\lambda$, was converted to intensity as a function of energy, f(E), in the energy range dE, using a Jacobian transformation:²⁷

$$f(E) = f(\lambda) \frac{dI_{\lambda}}{dE} \frac{d\lambda}{dE} = -f(I_{\lambda}) \frac{hc}{E^2}$$
 (S 7)

where h and c denote the Plank constant and the speed of light in vacuum, respectively.

The integrated areas (I_i) of the indicated transitions were calculated after removed the baseline and used the Jacobian transformation. The error in the integrated areas is estimated multiplying the standard deviation of noise by the integrated area, for each spectrum.

Thermometric performance of the Ln3+-containing micelles

Table S7. Parameters resulting from the fitting of a polynomial function $\triangle = A + BT + CT^2$ in the btfa- and DNPD-based polymeric micelles.

Ligand	A	B (10 ⁻³ K ⁻¹)	C (10 ⁻⁵ K ⁻²)	r ²
btfa	1.31±0.46	-8.96±0.30	1.59±0.47	0.988
DNPD	9.76±0.4	-6.51±0.26	11.38±0.42	0.982

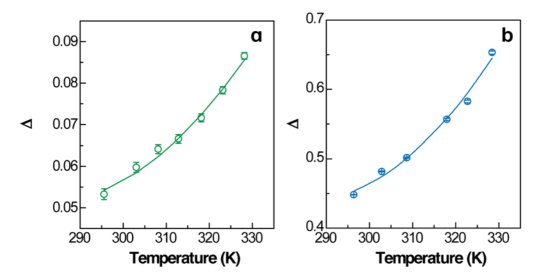


Figure S13. Calibration curve of **(a)** btfa- and **(b)** DNPD-based polymeric micelles. The lines are the fits to empirical functions with parameters presented in Table S7.

The relative thermal sensitivity (S_r) was calculated using:

$$S_r = \frac{1}{\Delta} \left| \frac{\partial \Delta}{\partial T} \right|. \tag{S 8}$$

and the temperature uncertainty, δT :

$$\delta T = \frac{\sqrt{2}}{S_r} \frac{\delta I(T)}{I(T)} \tag{S 9}$$

where $\delta I/I$ is a function of temperature.²⁸

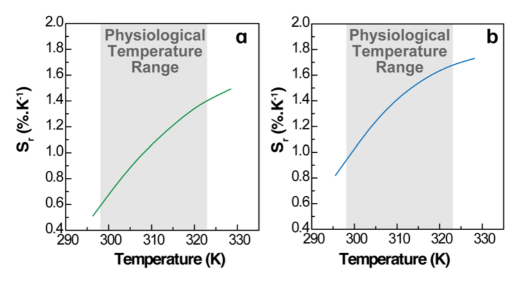


Figure S14. Relative thermal sensitivity, S_r , of **(a)** btfa- and **(b)** DNPD-based polymeric micelles. The physiological temperature range is shadowed.

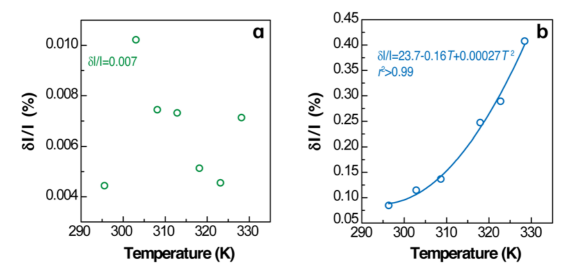


Figure S15. Signal-to-noise ratio, $\delta l/l$, in the calibration temperature range for **(a)** btfa-and **(b)** DNPD-based polymeric micelles. For DNPD-based material, a polynomial function was fitted to the experimental data (presented in (b)) whereas for the btfa-based material the arithmetic average value (0.007%) was used due to an erratic temperature dependence.

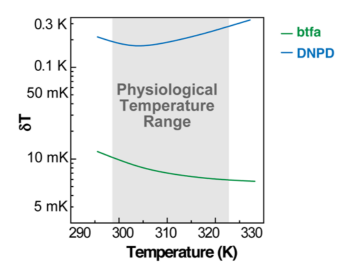


Figure S16. The temperature uncertainty, δT (T) of btfa- and DNPD-based polymeric micelles.

Cell cultures and temperature imaging

Cell cultures

MDA-MB-468 breast cancer cell lines were cultured in commercial DMEM (Dulbecco's Modified Eagle Medium) high glucose medium without phosphate (GIBCO), supplemented with 10% (v/v) fetal bovine serum (FBS), and 100 U/mL penicillin and 100 μg/mL streptomycin and kept in a thermostatic incubator (Heraeus Cell) in saturated humid air with 5% CO₂ at 37 °C (310 K). Successive subcultures and media changes were done every 2 or 3 days. Cells viability was calculated by the Trypan blue (Sigma) exclusion technique. Manipulations of cell cultures were carried out inside a vertical laminar flow cabinet (Telstar) in sterility conditions.

Preparation of cell samples for temperature imaging

In total, about 470,000 cells were seeded on coverslips in 6-well plates 24 h before the treatments with the micelles. Then, the cells were incubated with the DNPD-based polymeric micelles for 24 h, with different contents of the original suspension of the thermometric probes (0%, 10%, 20%, and 40%, v/v). In order to eliminate the presence of non-internalized nanoparticles, the supernatants were aspirated out and the cells were washed 3 times with cold NaCl 0.9%, fixed

with 4% (w/v) paraformaldehyde for 10 minutes, washed again three times with the same solution and mounted for microscopy. As a control, MDA-MB-468 cells were cultured in the absence of the Ln³⁺-bearing polymeric micelles and prepared in the same conditions.

The cell internalization ratio of the micelles was estimated by analysis of the lanthanide content in the supernatant after an incubation period of 24 h. The obtained values were 12%, 14% and 10% for the original suspension volume ratios in the culture of 10%, 20% and 40%, respectively.

Temperature imaging of fixed cells

Cell culture samples of cells incubated in the presence of the thermometric micelles were placed in a coverslip and examined in the microscope in three different imaging modes: (1) phase contrast, (2) full-color fluorescence, and (3) split fluorescent channels (for further temperature imaging). The cells cultures that have not incubated with the thermometric micelles were observed as a reference.

Temperature imaging of living cells

About 470 000 cells were cultivated in 35 mm glass-bottom dishes (ibidi) in DMEM in 2 ml of DMEM without phosphates with 10% SFB and antibiotics) for 24 h. Then, 0.2 mL of the thermometric micelle suspension was added and left incubating for another 24 h at 37°C. Then, we removed the medium, washed twice with DMEM medium without phosphates and SFB, and then two more times with 0.9% NaCl. Finally, we added 2 mL of 0.9% NaCl + glucose 0.9 mg·mL⁻¹ and the cell culture dishes were placed in a thermostatic holder as depicted in Figure S13. The temperature of the culture is measured with a fiber optic temperature sensor (Neoptix Reflex). The temperature imaging experiments were performed at different culture temperatures, 23, 28, 30 and 30.4°C, using an exposure time of 6s.

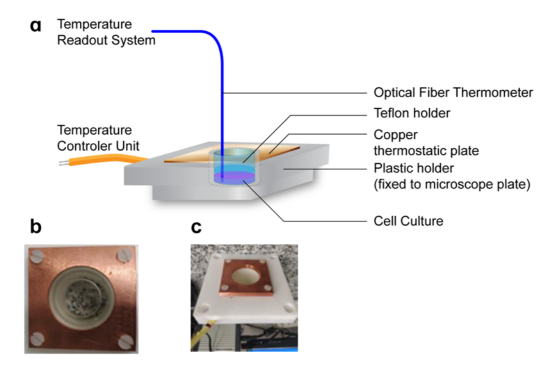


Figure S17. Thermostatic sample holder used in life cell imaging experiments. **(a)** General scheme of the sample holder **(b)** top and **(c)** perspective photograph of the copper thermostatic plate.

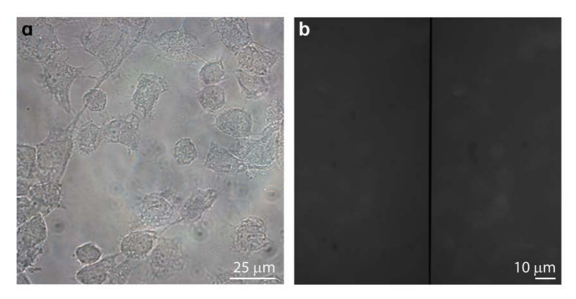


Figure S18. Control microscope images of control MDA-MB-468 cells before incubation with the micelles. **(a)** Phase contrast and **(b)** Split image using 365 nm illumination.

A drop of the DNPD-based polymeric micelles suspension was deposited on a 35 mm glass-bottom dishes, used previously for live-cells observation. The microscope image recorded in the Eu³⁺ channel (Figure S19a) shows a heterogeneous distribution of micelles in which the brighter spots (signed in yellow) correspond to high particle aggregation. We observe a narrow temperature distribution (fwhm~1 K), comparable to those recorded in the fixed MDA-MB-468 cells incubated with these micelles(Figure S19e).

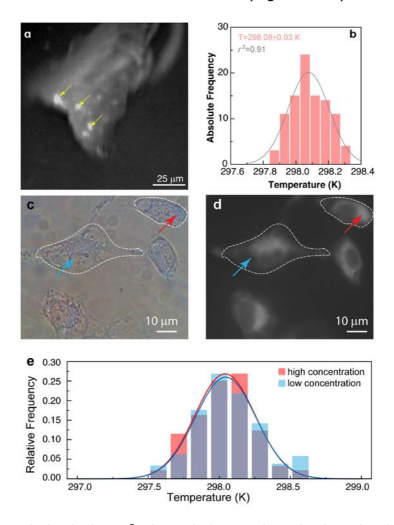
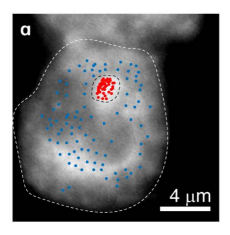


Figure S19. (a) Emission in the Eu³⁺ channel of DNPD-based polymeric micelles under UV-light illumination. The arrows mark the brighter spots. (b) Histogram of the temperatures measured in 100 points within the micelles. The solid line is the best fit to a Gaussian function (r^2 >0.91) with mean value±standard error of 298.08±0.03 K (the room temperature was measured as 298 K). (c) Phase contrast and (d) emission images in the Eu³⁺ channel of fixed MDA-MB-468 cells incubated with DNPD-based polymeric micelles under UV-light illumination. The blue and red arrows mark regions of high and low concentration of the thermometric probes, as can be observed in (c). (e) Temperature histograms on the regions signed with the blue and red arrows in (c) and (d). The lines are the best fits to experimental data using Gaussian functions (r^2 >0.97) corresponding to mean temperatures of 298.04±0.04 K and 298.04±0.05 K for the blue and red lines, respectively.



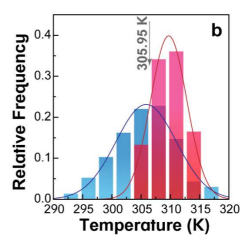


Figure S20. (a) Emission in the Eu³⁺ channel of MDA-MB-468 cells incubated with DNPD-based polymeric micelles recorded at 305.99 K. The blue and red points mark, respectively, the locations used for temperature determination of the darker (100 points) and brighter (50 points) regions. (b) Temperature histograms obtained from the selected points indicated in (a). The blue and red colors correspond to the blue (dark regions) and red (bright regions) points, respectively and the solid lines represent the best fits to Gaussian functions (r^2 >0.98).

The time dependence of the intracellular temperature while the external controller is setting the temperature of the culture was recorded (**Figure** S21). The temperature stabilization of the cells is monitored with a time step on the order of half-a-minute, and the steady-state regime is reached at about 3 minutes after the activation of the external temperature controller.

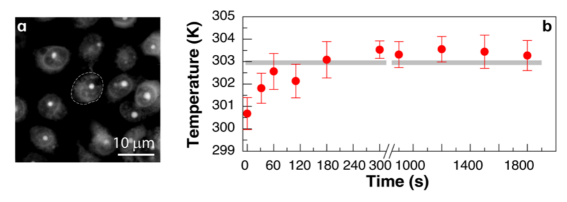


Figure S21. (a) Emission in the Eu³⁺ channel of MDA-MB-468 cells incubated with DNPD-based polymeric micelles under UV-light illumination (365 nm). The interrupted line delimits the cell whose temperature is tracked over time. (b) Temporal dependence of the mean temperature within the delimited cell. The error bars are the standard error and the horizontal bar marks the temperature set in the external temperature controller.

Optical properties of DNPD-based polymeric micelles in cell culture medium

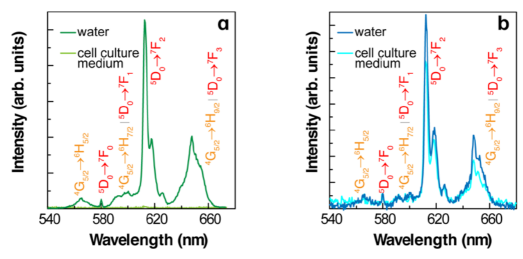


Figure S22. Room temperature emission spectra of (a) btfa- and (b) DNPD-based micelles under 355 nm and 405 nm excitation, respectively. The spectra were recorded for using water or DMEM cell culture media (incubated at 310 K for 24 h) as solvents.

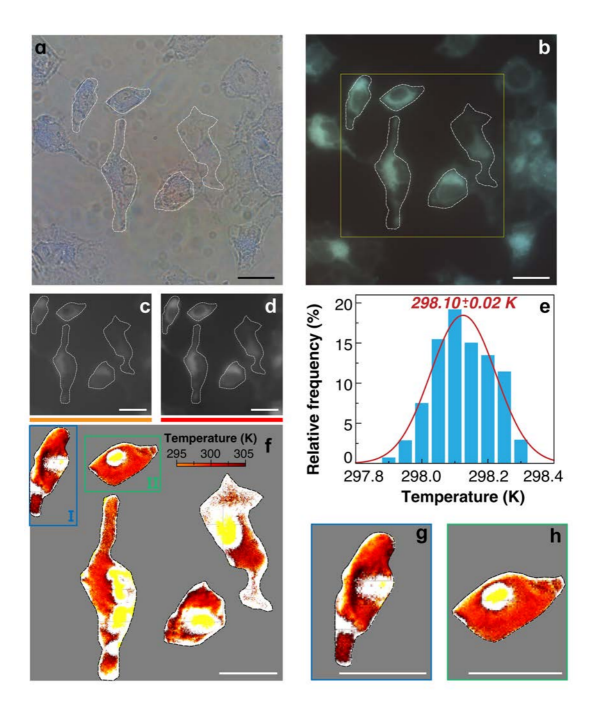


Figure S23. (a) Phase contrast and (b) fluorescence microscopy images of fixed MDA-MB-468 cells at 298.0 K. The yellow delimitated area in (b) is split in (c) I_1 and (d) I_2 . (e) The histogram of the temperature distribution using 100 points in the I and II areas follows a Gaussian curve of mean value±standard error 298.10±0.02 K (t^2 >0.95), accordingly with the cell culture temperature. The calculated temperature map of I_1 and I_2 channels is presented in (f). (g) and (h) Temperature mappings on the I and II areas, respectively. All scale bars correspond to 10 μm.

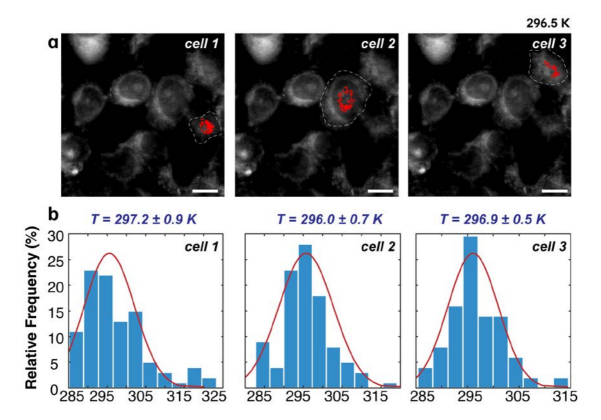


Figure S24. (a) Temperature maps of single living cells at 296.5 K. The temperature distributions were sampled in N>100 points (marked in red, randomly distributed in a radius of 3.25 μ m around the user-selected points) for each from 3 cells at indicated temperatures. Scale bars correspond to 10 μ m. (b) The corresponding temperature histograms constructed from the temperature distributions. The lines are the best fit to Gaussian functions ($r^2>0.89$) temperature values presented correspond to the mean±standard error.

Table S8. List of illustrative luminescent thermometers reporting intracellular temperature determination and temperature gradients (ΔT) within living cells.

Thermometer	Cell line	Location	Thermometric method	ΔT (K)	Ref.
QD655	HeLa, NIH3T3	Whole cell	Wavelength shift	~15	29
QD655	SH-SY5Y	Near the mitochondria	Intensity ratio	Large variance	30
Polymer encapsulated quantum dot	HepG2	Whole cell	Intensity	~15	31
Targeted organic dye	(HEK) 293	Mitochondria	Intensity	~15	32

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