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Eu³⁺ Sensitization via Nonradiative Interparticle Energy Transfer Using Inorganic Nanoparticles

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Cite This: J. Phys. Chem. Lett. 2020, 11, 689-695



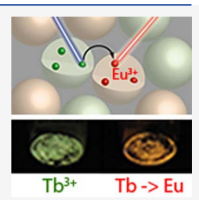
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ABSTRACT: Phosphors have been used successfully for both research and commercial applications for decades. Eu³⁺-doped materials are especially promising, because of their extremely stable, efficient, and narrow red emission lines. Although these emission properties are ideal for lighting applications, weak absorption in the blue spectral range has until now prevented the use of Eu³⁺-based phosphors in applications based on blue light-emitting diodes. Here, we demonstrate a sensitization mechanism of Eu³⁺ based on interparticle Förster resonance energy transfer (IFRET) between lanthanide-doped inorganic nanocrystals (NCs). Compared to co-doping different lanthanides in the same host crystal, IFRET allows an independent choice of host lattices for Eu³⁺ and its sensitizer while potentially greatly reducing metal-to-metal charge transfer quenching. We demonstrate IFRET between NCs, resulting in red Eu³⁺ emission upon blue excitation at 485 nm using LaPO₄:Tb/LaPO₄:Eu and LaPO₄:Tb/YVPO₄:Eu NC mixtures. These findings pave the way toward engineering blue-sensitized line emitters for solid-state lighting applications.



rivalent lanthanide-doped luminescent materials are of great interest for many applications because of their extremely stable and often narrow emission lines. They not only have been used successfully in fluorescent lighting applications, lasers, optical fibers, imaging, and displays for decades but also are candidates for sensing, labeling, and security applications. 1-5 Despite these successes, narrow-lineemitting trivalent lanthanides have limited suitability if excitation in the visible spectral range is desired, because their absorption is too weak outside the ultraviolet spectral range. For solid-state lighting applications, for example, narrow-line emitters such as Eu³⁺ would have ideal emission properties to boost color rendering and lumen efficacy.⁶ However, converter materials in solid-state lighting are usually pumped with a violet or blue light-emitting diode (LED), which has so far prevented these ions from being successfully applied in phosphors for white light LEDs.

A promising approach to increase the excitation probability of a lanthanide ion is by using a sensitizer, an optically active ion that strongly absorbs the wavelength of choice and then transfers the absorbed energy to the emitter ion (also called the activator) via nonradiative resonant energy transfer. For example, sensitizer ions like Eu²⁺, Ce³⁺, Pb²⁺, and Bi³⁺ have been considered to effectively enhance the absorption in the blue and sensitize Eu³⁺ emission.⁸⁻¹⁴ However, both the absorption properties of the sensitizer and the emission properties of Eu³⁺ depend on the host material. This severely limits the chances of finding a host material in which the sensitizer and Eu³⁺ can be co-doped and both have the desired properties. Another complication of co-doping is quenching of

the luminescence, which often occurs in nearest-neighbor pairs of sensitizers and emitters via a metal-to-metal charge transfer state (e.g., $Ce^{3+}-Eu^{3+} \rightarrow Ce^{4+}-Eu^{2+}$). This further narrows the selection of the emitter—sensitizer pair and their doping concentrations.

Charge transfer quenching can be circumvented by using energy transfer via energy migration over a concentrated sublattice of a third luminescent ion. This approach has been investigated in the past for Gd³⁺ and Tb³⁺ sublattices. 11,12,15 For example, the principle was reported for $Ce^{3+} \rightarrow (Tb^{3+})_n \rightarrow$ Eu^{3+} and $Eu^{2+} \rightarrow (Tb^{3+})_n \rightarrow Eu^{3+}$. In the latter system, increasing the Tb3+ concentration leads to more efficient energy migration, and above the percolation point of ~40% Tb³⁺, almost all of the migrating energy is captured by Eu³⁺ after excitation of Eu²⁺ at a wavelength (λ) of 377 nm. Although this approach works for high migration ion concentrations, metal-to-metal charge transfer quenching is easily thermally activated and still dominates at elevated temperatures.¹¹ In addition, energy migration effectively prevents charge transfer quenching only when no back-transfer to the sensitizer ion can occur, which is shown to be problematic for blue-excitable sensitizers such as cerium-doped garnets in combination with, for example, Tb3+ or Pr3+. 12,16

Received: December 18, 2019 Accepted: January 10, 2020 Published: January 10, 2020



Efficient nonradiative energy transfer and suppression of metal-to-metal charge transfer have also been demonstrated by physically separating sensitizer and emitter lanthanide ions in core/shell nanocrystal (NC) geometries ^{17–26} and have even been suggested for micrometer-sized particles. ²⁷ Core/shell geometries, however, greatly limit the choice of host lattice combinations because the crystal lattice parameters of the materials must be matched.

Here, we introduce a new approach for sensitizing Eu³⁺ emission that circumvents the issues of co-doping Eu³⁺ and a sensitizer into a single host lattice. It relies on interparticle Förster resonance energy transfer (IFRET) that couples nanocrystals (NCs) doped with sensitizer ions and NCs doped with Eu³⁺ (Figure 1). For efficient Förster energy

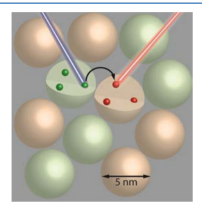


Figure 1. Sketch of the interparticle energy transfer mechanism. Red light is emitted from the Eu³⁺-doped emitter particle, following excitation of a neighboring nanoparticle doped with Tb³⁺ sensitizer ions. Due to the physical separation of both emitter and sensitizer ions, charge transfer quenching is prevented.

transfer between lanthanide ions, sensitizer—emitter distances on the order of 0.5–1.5 nm are required. ^{28,29} The concept of IFRET in NC composites provides various tuning knobs to control the distances and transfer efficiencies between sensitizer and emitter ions (see Figure S1). To achieve successful IFRET, we therefore ensure that the NCs in our multihost-lattice composite material are small and in the proximity of each other. Our approach allows an independent choice of the sensitizer and emitter host lattice and prevents charge transfer quenching, which is an even shorter-distance process (<0.5 nm range) than energy transfer via dipolar interactions (0.5–1.5 nm). 11,30,31 IFRET has previously been demonstrated between lanthanide ions only in the same host materials and never with materials that were blue-excitable. 32,33 In this work, we demonstrate efficient sensitization of Eu³⁺ via IFRET from Tb³⁺ ions in different material systems. We show IFRET sensitization in the blue at a λ of 485 nm and achieve IFRET even if the host materials for emitter and sensitizer ions are different. We also demonstrate the robustness of the IFRET-sensitized Eu³⁺ emission intensity from room temperature to 145 °C.

We first tested the homogeneous mixing of the Eu³⁺-doped emitting NCs and the Tb³⁺-doped sensitizer NCs. To this end, "wet-mixed" NC composites were compared with "dry-mixed" composites (see Experimental Methods for details). Scanning transmission electron microscopy (STEM) combined with energy dispersive X-ray spectroscopy (EDX) elemental mapping (Figure S2) shows that the NCs are approximately

5–10 nm in diameter with doping concentrations of 10% Eu³⁺ in YVPO₄, 40% Tb³⁺ in LaPO₄, and 7% Eu³⁺ in LaPO₄. High-resolution EDX mappings on LaPO₄:Tb/YVPO₄:Eu NC composites confirm that the wet-mixed sample is a homogeneous composite of the two types of NCs (Figure 2B,D), showing strong intermixing of the two materials on a

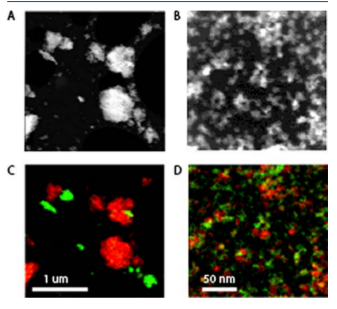


Figure 2. (A and B) HAADF-STEM images and (C and D) STEM-EDX mappings of (A and C) dry-mixed and (B and D) wet-mixed LaPO₄:Tb/YVPO₄:Eu NCs. Green represents lanthanum (in LaPO₄:Tb), and red represents vanadium (in YVPO₄:Eu). Clearly, dry mixing results in inhomogeneities at micrometer length scales, while wet mixing produces homogeneous composites.

<10 nm scale. The dry-mixed sample, on the other hand, is largely inhomogeneous (Figure 2A,C), with very little intermixing between the two types of NCs on a scale of less than hundreds of nanometers. More STEM-EDX mappings of these samples are shown in Figure S3. Because FRET between lanthanide ions typically works effectively only for donoracceptor distances of <1.5 nm, NC intermixing on a scale of a few hundred nanometers is not expected to result in any measurable IFRET signals. We therefore use the dry-mixed sample as a reference without IFRET. The same structural analysis could not be performed on the LaPO₄:Tb/LaPO₄:Eu mixtures, because it proved to be too difficult to distinguish between these materials using EDX. Nevertheless, our optical measurements (see below) confirm similar structural differences of mixing between wet-mixed and dry-mixed samples for the LaPO₄:Tb/LaPO₄:Eu composites.

Next, the samples were characterized optically to obtain evidence for interparticle energy transfer. In the case of IFRET, excitation in the 5D_4 level of Tb^{3+} will result in red emission from the 5D_0 level of Eu^{3+} and give rise to extra Eu^{3+} emission lines, which serves as a fingerprint for IFRET. In excitation spectra of a characteristic Eu^{3+} emission line, the Tb^{3+} 7F_6 - 5D_4 excitation line will appear as a result of IFRET. Figure 3 shows the emission spectra of the wet-mixed and dry-mixed $LaPO_4$: $Tb/LaPO_4$:Eu NC composites upon 5D_4 excitation of Tb^{3+} at a λ of 484.5 nm. As a comparison, we also show the emission of the single-component NC composites, $LaPO_4$:Tb and $LaPO_4$:Eu. The single-component $LaPO_4$:Eu NCs show strong emission lines centered at λ values of 540, 590, and 620

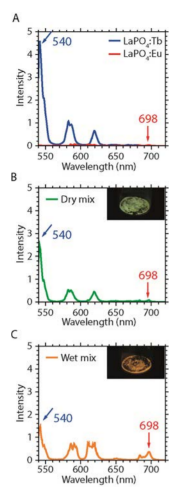


Figure 3. Emission spectra upon Tb³+ excitation at a λ of 484.5 nm of (A) single-component LaPO₄:Tb (blue) and LaPO₄:Eu (red) NC composites and (B) dry-mixed and (C) wet-mixed LaPO₄:Tb/LaPO₄:Eu NC composites. A Eu³+ emission band is observed at a λ of 698 nm in the case of the wet-mixed NCs, indicating Eu³+ emission upon Tb³+ excitation. The insets in panels B and C show photographs of the luminescence of the dry- and wet-mixed materials upon excitation at 365 nm.

nm, corresponding to the ${}^5D_4 \rightarrow {}^7F_5$, ${}^5D_4 \rightarrow {}^7F_4$, and ${}^5D_4 \rightarrow {}^7F_3$ transitions of Tb³⁺, respectively. The single-component LaPO₄:Eu NCs have almost no luminescence over the entire range when excited at a λ of 484.5 nm as there is no resonant absorption line of Eu³⁺ at this wavelength.

The dry-mixed NC composite shows a reduced $Tb^{3+} {}^5D_4 \rightarrow$ ⁷F₅ emission intensity around 540 nm (Figure 3B) compared to the single-component LaPO₄:Tb NCs (Figure 3A). This results from dilution as a result of the 1:1 weight ratio mixing with LaPO₄:Eu NCs. For the wet-mixed sample (Figure 3C), the 540 nm emission line is even weaker than for the drymixed sample (Figure 3B). In addition, we observe the appearance of new emission lines around 590, 610, and 698 nm that can be ascribed to ${}^5D_0 \rightarrow {}^7F_1$, ${}^5D_0 \rightarrow {}^7F_2$, and ${}^5D_0 \rightarrow$ ⁷F₄ emission of Eu³⁺, respectively. The 590 and 610 nm emissions overlap strongly with Tb3+ emission lines, but the 698 nm Eu³⁺ emission is well-separated from Tb³⁺ emission and can serve as a characteristic Eu³⁺ emission line. These observations suggest energy transfer from Tb3+ to Eu3+ in the wet-mixed NC composite, while the dry-mixed sample does not show indications of IFRET. The differences in IFRET

efficiencies for wet- and dry-mixed samples are consistent with our expectations based on structural analysis (Figure 2). This highlights the importance of homogeneous intermixing at the nanometer scale for efficient IFRET. The photographs in the insets of Figure 3 show that the luminescence color difference between the two samples is clearly visible by eye, where the wet-mixed NCs clearly emit more red (Eu³⁺) light, compared to the dry-mixed NCs that mostly show green (Tb³⁺) emission.

To confirm preservation of the NC quality during the mixing procedures, the Eu $^{3+}$ emission was measured for direct excitation ($^7F_0 \rightarrow ^5L_6$) at 395 nm (Figure S4). The Eu $^{3+}$ emission spectrum and intensity are very similar for the wetand dry-mixed samples, indicating similar mixing ratios and preservation of the LaPO₄:Eu NC integrity in the two mixed samples.

Additional evidence for IFRET is found by measuring the excitation spectra of the Eu³+ $^5D_0 \rightarrow ^7F_4$ emission peak at a $\lambda_{\rm em}$ of 698 nm for both mixtures and for the individual single-component materials (for the pure LaPO4:Tb sample, we monitored at a $\lambda_{\rm em}$ of 542 nm), as shown in Figure 4. The excitation spectra of dry-mixed NCs and single-component LaPO4:Eu are identical (Figure 4B), confirming the absence of Eu³+ sensitization by Tb³+. In contrast, the excitation spectrum of the wet-mixed NCs shows clear evidence of Tb³+ excitation lines (most notably, between 340 and 380 nm and at 485 nm), in addition to those originating from Eu³+ (most notably, at 395 and 465 nm). This is consistent with IFRET in the wet-mixed sample.

IFRET is a nonradiative decay pathway for the Tb^{3+} 5D_4 excited state and will result in faster and non-exponential decay of the 5D_4 emission. Measuring luminescence decay curves provides additional evidence for IFRET and helps to quantify the average transfer rate and efficiency. Figure 5 shows decay curves of Tb^{3+} 5D_4 emission ($\lambda_{\rm em}=542$ nm). The average lifetimes for the dry- and wet-mixed samples are 1.5 and 0.7 ms, respectively (1/e decay time). The decrease in decay time of the donor emission is a well-known signature for FRET. On the basis of these numbers, we can estimate that the average IFRET rate in our wet-mixed sample is ~ 0.76 ms⁻¹. Assuming equal nonradiative decay rates from other mechanisms, we can estimate the IFRET efficiency in the wet-mixed sample from the ratio between the integrated areas under the decay curves³⁵ to be 54%.

To investigate the temperature stability of the IFRETsensitized emission, emission spectra were recorded upon Tb³⁺ excitation at 485 nm from room temperature to 145 °C (Figure 6). Interestingly, the Eu³⁺ emission stays almost constant within this temperature range. The Tb3+ emission shows some thermal quenching behavior. The intensity recovers when the sample is cooled again (Figure S5b). Temperature-dependent excitation spectra (Figure S5c,d) do show some thermal quenching of the Eu3+ emission upon direct excitation (e.g., at $\lambda = 395$ nm), but not for sensitization via IFRET, which can be explained by a slightly higher IFRET efficiency at elevated temperatures. An incidental increased Eu³⁺ emission intensity is observed at 105 °C but seems to be due to an experimental error, because it is not observed in the "back trace" (see Figure S5b). In addition, the excitation spectra measured at 105 °C (Figure S5c) show enhanced Eu³⁺ emission for not only excitation via energy transfer (i.e., at λ = 485 nm) but also direct excitation (i.e., $\lambda = 395$ nm), indicating that the increased Eu³⁺ emission at 105 °C is not the result of enhanced energy transfer efficiencies. For the sake of

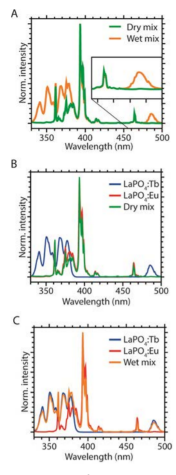


Figure 4. Excitation spectra of $\rm Eu^{3+}$ emission, monitored at a λ of 698 nm, of (A) dry-mixed (green) and wet-mixed $\rm LaPO_4$:Tb/LaPO₄:Eu (orange) NCs, (B) dry-mixed (green) NCs, together with pure $\rm LaPO_4$:Tb (blue) and $\rm LaPO_4$:Eu (red) NCs, and (C) wet-mixed (orange) NCs, together with pure $\rm LaPO_4$:Tb (blue) and $\rm LaPO_4$:Eu (red) NCs. The inset in panel A shows a close-up of the area between 450 and 500 nm. Clear differences are observed between the wet-mixed and dry-mixed NC excitation spectra. For the dry-mixed NCs, a typical $\rm LaPO_4$:Eu excitation spectrum is observed, whereas both $\rm LaPO_4$:Eu and $\rm LaPO_4$:Tb excitation lines are observed in the case of wet-mixed NCs.

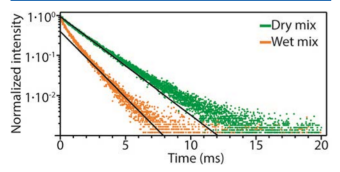


Figure 5. Photoluminescence decay curves of the Tb^{3+} $^5\mathrm{D}_4$ state in dry-mixed and wet-mixed LaPO₄:Tb/LaPO₄:Eu NC composites. Black lines represent single-exponential fits, using decay times of 1.77 and 1.32 ms for the dry and wet mix, respectively. Excitation was at a λ of 355 nm, emission was monitored at a λ of 542 nm.

completeness, photoluminescence decay curves at elevated temperatures are shown in Figure S6. The temperature robustness of sensitization via IFRET of both emission

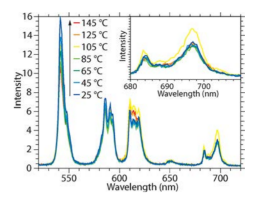


Figure 6. Temperature-dependent emission spectra on LaPO₄:Tb/LaPO₄:Eu wet-mixed NCs. The inset shows a close-up of the Eu³⁺ emission between 680 and 720 nm, which is almost constant over the measured temperature range. The sample was excited in the Tb³⁺ excitation line at a λ of 484.5 nm.

intensity and peak wavelength is a significant advantage of lanthanide-based multihost composites over, e.g., quantum dots or perovskite emitter materials for lighting applications.

To investigate IFRET between the sensitizer and emitter incorporated in different host lattices, we investigated the emission and excitation spectra of wet- and dry-mixed NC composites of LaPO₄:Tb and Y(V₁P)O₄:Eu (Figure 7). Similar to what we observed for the LaPO₄:Tb/LaPO₄:Eu mixtures, only the wet-mixed sample shows Eu³⁺ emission at 698 nm upon Tb³⁺ excitation at 485 nm (Figure 7A,B). Many of the Tb³⁺ excitation lines are overlapping with the strong vanadate charge transfer band starting at 370 nm, but the Tb³⁺ excitation peak at 485 nm is clearly observed in the excitation spectrum of 698 nm Eu³⁺ emission for the wet-mixed sample but not for the dry-mixed sample (Figure 7D-F). Emission spectra for direct Eu³⁺ excitation at 395 nm are shown in Figure S7. Photoluminescence decay curves of the Tb^{3+ 5}D₄ emission (Figure 7C) confirm nonradiative energy transfer in the wetmixed sample. Taking the 1/e decay time leads to estimates of 9.2 ms⁻¹ for the average IFRET rate and 78% for the IFRET efficiency. The oscillator strength of the acceptor and distance distribution affect the IFRET rate and are dependent on the host lattice of the Eu³⁺ ions. This could explain the difference between IFRET rates for LaPO4:Tb/LaPO4:Eu and La-PO₄:Tb/YVPO₄:Eu.

The observations depicted in Figure 7 provide evidence for efficient nonradiative energy transfer from ${\rm Tb}^{3+}$ in ${\rm LaPO_4}$ to ${\rm Eu}^{3+}$ in ${\rm YVPO_4}$, thereby demonstrating IFRET between dopants in different host materials for the first time. Even though IFRET efficiencies are generally lower than FRET efficiencies that can be observed for intraparticle energy transfer, near-unity efficiencies are not needed for many applications. To use ${\rm Eu}^{3+}$ as a red emitter in white LEDs, for example, IFRET efficiencies of only 20–30% are high enough to achieve high color rendering warm white light when a green emitter (such as ${\rm LaPO_4:Tb}$ or ${\rm Y_3Al_5O_{12}:Ce}$) is used as the sensitizer (Figure S8). The reported efficiencies of 54% and 78% for ${\rm LaPO4:Tb} \rightarrow {\rm LaPO4:Eu}$ and ${\rm LaPO4:Tb} \rightarrow {\rm YVPO4:Eu}$, respectively, are thus high enough to be interesting for these kinds of applications.

In conclusion, we have demonstrated IFRET between lanthanide-doped NCs. For LaPO₄:Tb/LaPO₄:Eu and La-PO₄:Tb/YVPO₄:Eu nanocrystals homogeneously intermixed at the nanoscale, we observed efficient sensitization of Eu³⁺

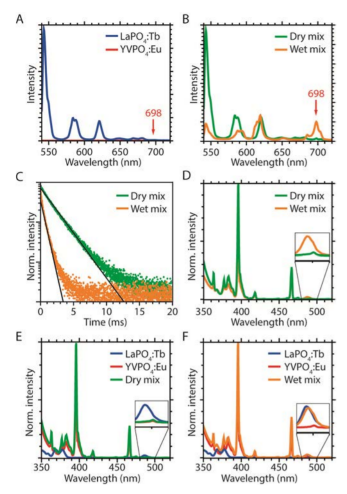


Figure 7. Photoluminescence measurements on LaPO₄:Tb/YV-PO₄:Eu NCs. (A and B) Emission spectra upon Tb³⁺ excitation at a λ of 484.5 nm of (A) single-component LaPO₄:Tb (blue) and YVPO₄:Eu (red) NCs and (B) dry-mixed (green) and wet-mixed (orange) LaPO₄:Tb/YVPO₄:Eu NC composites. (C) Luminescence decay curves of the Tb3+ 5D4 emission in dry-mixed (green) and wetmixed (orange) LaPO₄:Tb/YVPO₄:Eu NC composites. Excitation at a λ of 355 nm, emission was monitored at a λ of 542 nm. Black lines represent single-exponential fits, using decay times of 1.95 and 0.61 ms for the dry and wet mix, respectively. (D-F) Excitation spectra of Eu³⁺ emission, monitored at a λ of 698 nm, of (D) dry-mixed (green) and wet-mixed (orange) LaPO₄:Tb/YVPO₄:Eu NC composites, (E) dry-mixed (green) NC composites, together with single-component LaPO₄:Tb (blue) and YVPO₄:Eu (red) NCs, and (F) wet-mixed (orange) NCs, together with single-component LaPO4:Tb (blue) and YVPO₄:Eu (red) NCs. The insets show a close-up of the area between 480 and 500 nm.

emission upon Tb³⁺ excitation at 484.5 nm. Further support for IFRET is provided by excitation spectra of Eu³⁺ emission that clearly show the 485 nm Tb³⁺ excitation line and trace the full Tb³⁺ excitation spectrum in addition to the expected Eu³⁺ excitation lines. Furthermore, luminescence decay curves acceleration of the Tb³⁺ ⁵D₄ emission in intermixed material, confirming nonradiative energy transfer from Tb³⁺ to Eu³⁺ acceptor ions in neighboring NCs. We envision that the concept of IFRET can be applied for a wide variety of sensitizer and emitter ions and in a wide variety of nanocrystal host materials. The flexibility in choice of host lattice for the sensitizer and emitter gives unprecedented freedom in independently tuning excitation and emission properties and

opens up a whole new world of engineering of luminescent materials. For example, in new nanocomposites, efficient lanthanide emission can be generated upon visible light excitation, paving the way toward using lanthanides for a myriad of solid-state lighting applications.

EXPERIMENTAL METHODS

Materials. Lanthanum(III) chloride hexahydrate (LaCl₃·6H₂O, 99.9% La, Brunschwig Chemie BV), europium(III) chloride hexahydrate (EuCl₃·6H₂O, 99.9% trace metal basis, Sigma-Aldrich), terbium(III) chloride hexahydrate (TbCl₃·6H₂O, 99.9% trace metal basis, Sigma-Aldrich), methyl alcohol (methanol, anhydrous, 99.8%, Sigma-Aldrich), tributyl phosphate (≥99.0%, Sigma-Aldrich), diphenyl ether (99%, Sigma-Aldrich), tributylamine (≥99.0%, Sigma-Aldrich), phosphoric acid (≥99.999%, Sigma-Aldrich), dihexyl ether (97%, Sigma-Aldrich), toluene (anhydrous, 99.8%, Sigma-Aldrich), an ammonium hydroxide solution (30%, Sigma-Aldrich), hydrogen peroxide (30 wt % in H₂O, Sigma-Aldrich), ethyl alcohol (ethanol, 30 ppm of H₂O, VWR international), hydrochloric acid (HCl, 37% in water, Sigma-Aldrich), and mixed phosphate/vanadate Y(V,P)O₄:Eu (YVPO4:Eu) nanoparticles (7% Eu, CAN GmbH, CANdot Series X) were used. HCl was diluted with water to 0.1 M, and all other chemicals were used as received.

Sample Preparation. LaPO₄:Eu³⁺ (10%) and LaPO₄:Tb³⁺ (40%) NPs with tributylamine ligands were synthesized using the procedure reported by the group of Haase.³⁶⁻³⁹ In the case of LaPO4:Eu NPs, 9 mmol of LaCl₃·6H₂O and 1 mmol of EuCl₃·6H₂O were used, whereas for LaPO4:Tb, 6 mmol of LaCl₃·6H₂O and 4 mmol of TbCl₃·6H₂O were dissolved in approximately 10 mL of methanol until a clear solution was acquired. Then, 40 mmol of tributyl phosphate was added, and methanol was removed from the reaction mixture under vacuum using a Schlenk line. Thirty milliliters of diphenyl ether was added before the system was flushed with N₂. Next, water was released under vacuum at 105 °C before the reaction mixture was cooled to <50 °C and 40 mmol of tributylamine was added. Finally, 7.0 mL of a 2 M phosphoric acid solution in dihexyl ether was added. The reaction mixture was heated to 200 °C and left to react for 16 h before being cooled back to room temperature. The nanocrystals were collected by centrifuging and washed several times with toluene. The obtained nanoparticles were dried for further treatment and analysis. X-ray diffraction (XRD) patterns were measured for all nanocrystals to confirm the crystal structures before using them in the mixtures (Figure S9).

For the "wet-mixed" NC composites, 200 mg of dried LaPO₄:Eu or YVPO₄:Eu nanoparticles and 200 mg of dried LaPO₄:Tb nanoparticles were mixed in a vial and dispersed in 20 mL of water. The mixing was enhanced by sonication in an ultrasonic bath for 90 min. For the "dry-mixed" NC composites, 100 mg of dried LaPO₄:Eu nanoparticles and 100 mg of dried LaPO₄:Tb nanoparticles were separately dispersed in 20 mL of water or ethanol in different vials.

Ligands of the LaPO₄ nanoparticles were removed using a base-piranha treatment. The piranha solution was prepared by mixing an \sim 3:1 weight ratio of a NH₄OH solution (30% in water, heated to 60 °C) and H₂O₂ (30% in water); 20–50 mL of this solution was added to the nanoparticle dispersions. The mixture was left to react at 70–80 °C for 90 min before the mixture was cooled to room temperature. The nanoparticles were re-collected by centrifugation and washed multiple times

with acidic ethanol (0.1 M HCl was added to ethanol to obtain a pH of 4).

The NC dispersions were dried at 120–150 °C in the oven. The mixed dispersion thus formed a wet-mixed NC composite. The dry NCs from the single-component dispersions were mixed with a mortar and pestle after drying to form dry-mixed NC composites, as reference samples.

Characterization. High-resolution transmission electron microscopy (HRTEM) images were taken using a JEOL ARM 200F instrument. The transmission electron microscope was probe corrected, equipped with a 100 mm² Centurio SDD EDX detector, and operated at 200 kV. Samples were prepared by dispersing the dry powder in ethanol, followed by sonication for 2 min. The samples were drop casted on a copper grid with holey carbon film.

Emission and excitation spectra were recorded using an Edinburgh Instruments FLS928 spectrofluorometer. A 450 W xenon lamp in combination with a monochromator was used for excitation, while the emitted light was collected using a R928 photomultiplier tube ($\lambda = 400-850$ nm). To record luminescence decay curves, the samples were excited using an optical parametric oscillator (OPO) system (Opotek Opolette HE 355 II), pumped by the third harmonic (355 nm) of a YAG:Nd laser, with a pulse width of 5 ns and a repetition rate of 20 Hz. For temperature-dependent measurements, a LINKAM cell was used, which has an optical window to monitor emission and excitation spectra during heating. All optical measurements were performed in reflection mode on optically thick powder samples.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpclett.9b03764.

Supporting simulations, XRD data, and additional (S)TEM, (S)TEM-EDX, and optical data (PDF)

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Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

Marcel Verheijen from Eurofins Material Science Netherlands BV is gratefully acknowledged for technical support for HRTEM imaging and elemental mapping. Pradip Chakraborty and Jacobine van Hest are thanked for providing LaPO₄ nanoparticles, and Valerio Favale is thanked for photographs of the samples. This work is part of the research program Innovation Fund Chemistry (LIFT) with Project 731.017.401, which is (partly) financed by the Dutch Research Council (NWO). F.T.R. is supported by NWO Veni Grant 722.017.002 and by The Netherlands Center for Multiscale Catalytic Energy Conversion (MCEC), an NWO Gravitation program funded by the Ministry of Education, Culture and Science of the government of The Netherlands.

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