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Interplay between Static and Dynamic Energy Transfer in Biofunctional Upconversion Nanoplatforms

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Supporting Information

ABSTRACT: Clarification of the energy-transfer (ET) mechanism is of vital importance for constructing efficient upconversion nanoplatforms for biological/biomedical applications. Yet, most strategies of optimizing these nanoplatforms were casually based on a dynamic ET assumption. In this work, we have modeled quantitatively the shell-thickness-dependent interplay between dynamic and static ET in nanosystems and validated the model in a typical biofunctional upconversion nanoplatform composed of NaYF₄:Er, Yb/NaYF₄ upconversion nanoparticles (UCNPs), and energy-acceptor photosensitizing molecule Rose Bengal (RB). It was determined that with a proper thickness shell, the energy transferred via dynamic ET as well as static ET in this case could be significantly improved by ~4 and ~9 fold, respectively, compared with the total energy transferred from bare core UCNPs. Our results shall form the bedrock in designing highly efficient ET-based biofunctional nanoplatforms.

Lanthanide ion (Ln³⁺)-doped upconversion nanoparticles (UCNPs) able to convert continuous-wave near-infrared (NIR) light into higher-energy and multicolor UV/visible light have been exploited intensively for various biological and biomedical applications.¹−³ They are more attractive than currently used materials in, for example, deep tissue penetration, and high detection sensitivity ascribed to the low photobleaching and absence of fluorescence interference of the background biological environment. Most of these applications are based on different energy-transfer (ET) mechanisms, for example, biosensing asks for high dynamic ET ( Förster or fluorescence resonant energy transfer (FRET)) or static ET (inner filter effect or reabsorption), whereas photodynamic/thermal therapy requires high total ET including static and dynamic ET.⁴−⁶

In 2005, a homogeneous biosensor was reported using UCNPs as the energy donor.¹⁰ In the presence of avidin, biotin conjugated Na(Y₁₋ₓNaₓ)F₄:Yb³⁺, Er³⁺/Tm³⁺ UCNPs (∼50 nm) and Au-biotin nanoparticles (the acceptor) were brought into close proximity through specific interaction between the avidin and biotin, and the upconversion luminescence was quenched via dynamic ET. On the basis of this effect, simple and sensitive detection of trace amounts of avidin was realized. Since then, many other nanomaterials and organic dyes, such as carbon nanoparticles, graphene oxide, and tetramethylrhodamine, have been introduced as energy acceptors into UCNPs-based biochemical analysis of proteins, DNA, heavy metals, enzyme activities, and so forth.¹,¹¹−¹⁴ The static ET process, where close proximity between the donor and acceptor is not required, was also utilized for the sensing of pH, CO₂, Cr⁶⁺, and so forth.¹⁵−¹⁸ In the meantime, other ET-based applications, for example, photodynamic therapy employing photosensitizing molecule conjugated upconversion nanoplatforms, were introduced as well.¹⁸−²⁰

However, to bring these proof-of-concepts to application, there are huge challenges. One crucial issue is that the ET needs to be largely enhanced because the state-of-the-art upconversion efficiency is below 1% for nanomaterials under a clinically acceptable excitation level.¹⁹ Until now, the picture of the ET mechanism has not been well explored, as witnessed by

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the fact that most relevant ET has been treated as dynamic in nature. The distance dependence of FRET was generally taken as the basis for optimizing the biofunctional nanoplatforms that led to the popular approach (shortening the distance between the UCNPs and the energy acceptors), and naturally, a bare-core structure was dominantly adopted.\textsuperscript{1,\textsuperscript{2}2\textsuperscript{2},2\textsuperscript{6}–2\textsuperscript{9}} Recently, there were some indications that ET mechanisms other than FRET might not be neglected between a closely located donor and acceptor.\textsuperscript{27} For example, reabsorption was shown to be the primary reason for the luminescence quenching of NaYF\textsubscript{4}:Yb\textsuperscript{3+}, Er\textsuperscript{3+} UCNPs (\(\sim 23.5\) nm, the donor) encapsulated by an amphiphilic polymer shell containing photosensitive diarylethene chromophores (the acceptor).\textsuperscript{30} Similar phenomena were also reported when CdSe quantum dots were used as the energy acceptor in the solid state.\textsuperscript{31}

Therefore, to have a quantitative picture of the roles that the static and dynamic ET play in the biofunctional nanoplatforms is very urgent for constructing efficient ET structures and of key importance for bioapplications of these very promising nanoplatforms. In this work, we separate quantitatively the dynamic and static ET and their weights in ET from theoretical modeling and corresponding spectroscopic experiments and validate the picture using a typical biofunctional nanoplatform. In a dynamic ET process, energy is transferred to the acceptor nonradiatively from the excited state of the donor, providing an additional nonradiative relaxation channel for the donor emissive energy level, resulting in a shortened lifetime of the donor emission. Whereas in the static ET process the energy is transferred to the acceptor after the radiative relaxation process of the donor, the temporal behavior of the donor emission is thus not affected.

Considering the spatial distribution of the emitting centers and the relatively large size of UCNPs (compared with typical Förster distance of less than 10 nm), only the emitting centers close to the surface-bound acceptors can transfer energy effectively via dynamic ET. Therefore, we can reasonably divide the doped area of UCNPs into two parts; one is the outer layer where the emitting centers participate in both dynamic and static ET and thus have fast emission kinetics, and the other one is the central area where the emitting centers are farther away from surface-bound acceptors and thus not involved in dynamic ET, retaining slow emission kinetics.

An important biofunctional upconversion nanoplatform is taken as the sample system in this case, which is composed of core–shell NaYF\textsubscript{4}:Yb\textsuperscript{3+}, Er\textsuperscript{3+}/NaYF\textsubscript{4} and a covalently conjugated photosensitizing molecule, Rose Bengal (RB) (Figure 1A), and the absorption and emission spectra of NaYF\textsubscript{4}:Yb\textsuperscript{3+}, Er\textsuperscript{3+} UCNPs and RB are depicted in Figure 1B. Obviously, NIR 980 nm excitation, once upconverted in UCNPs to \(\sim 540\) nm, can be transferred to RB via static and/or dynamic ET mechanisms due to the large overlap of the 540 nm (green) emission band and the absorption spectrum of RB, whereas the upconversion emission band at \(\sim 650\) nm (red) does not participate in the ET because it does not overlap spectrally with RB absorption.

Assuming that the time evolution of the green emission of UCNPs without the acceptor RB under NIR light excitation is described with a biexponential function, that is

\[
I(t)_{\text{D}} = A_1 \exp \left( -\frac{t}{\tau_{\text{rise},\text{RB}}} \right) + A_2 \exp \left( -\frac{t}{\tau_2} \right)
\]

(1)

where \(\tau_2\) is the decay time constant. When bounded on the surface of the UCNPs, RB can be excited (1) via static ET, that is, absorption of the 540 nm emission of UCNPs, and (2) via dynamic ET. Therefore, the time evolution of the 540 nm upconversion emission can be approximately described as

\[
I(t)_{\text{D-Â}} = A_1 \exp \left( -\frac{t}{\tau_{\text{rise},\text{RB}}} \right) + A_2 \exp \left( -\frac{t}{\tau_2} \right) + A_3 \exp \left( -\frac{t}{\tau_3} \right)
\]

(2)

Here, the additional short decay component with time constant \(\tau_3\) represents emitters in the outer layer of UCNPs that experience dynamic ET. \(\tau_{\text{rise},\text{RB}}\) represents the rise time constant when RB exists.

Because the lifetime of upconversion luminescence (micro- to milliseconds) is much longer than that of RB (pico- to nanoseconds), the time evolution of RB emission shall mimic that of upconversion luminescence when they are excited via ET from UCNPs regardless of being dynamic or static.\textsuperscript{32–34} Therefore, the temporal behavior of RB emission can also be described with a triexponential function with the time constants of eq 2.
The amplitude ratio $A_2/A_3$ of the slow to the fast decay components in eq 2 represents also the ratio of the numbers of emitting centers in the central area to the outer layer of UCNPs. The amplitude ratio $A_2'/A_3'$ provides more important information about acceptor excitation. It is known that dynamic ET efficiency depends on the donor–acceptor distance with an inverse $n$th power law ($n > 3$), while the static ET rate drops off according to the inverse square law in relation with the absorption cross section of the acceptor; static ET is thus less sensitive to the separation distance. On the basis of this, we can separate the contributions of dynamic ET and static ET of the emitters in the outer layer of UCNPs by rewriting eq 3 to

$$I'(t)_{D-D} = A_1' \exp\left(-\frac{t}{\tau_{\text{rise, RB}}}\right) + A_2' \exp\left(-\frac{t}{\tau_2}\right) + A_3' \exp\left(\frac{t}{\tau_3}\right)$$

(3)

where $A_1' = A_2' + B$, $A_2'$ denotes the contribution of static ET, and $B$ denotes the contribution of dynamic ET. Because

$$\frac{A_2'}{A_3'} = \frac{A_2}{A_3}$$

(5)

the ratio between the contributions of dynamic and static ET from the emitting centers in the outer layer area is thus

$$D = \frac{B}{A_2'} = \frac{A_2A_3'}{A_2'A_3} - 1$$

(6)

To validate this model, we have varied the shell thickness of the UCNPs and monitored the time evolution of the upconversion emission and RB emission. NaYF$_4$:Yb$^{3+}$, Er$^{3+}$ core, and NaYF$_4$:Yb$^{3+}$, Er$^{3+}$/NaYF$_4$ core–shell UCNPs with increasing shell thickness (from cs1 to cs5) were synthesized following a self-focusing strategy utilizing Ostwald ripening dynamics. XRD patterns are shown in Figure S1 in the Supporting Information (SI), confirming the hexagonal phase of all of the samples. Figure 2 illustrates the high monodispersity of the samples.

Steady-state emission spectra of the six samples were recorded, as shown in Figure 3. Despite the continuous increase of the green and red upconversion emission and monotonous decrease of ET efficiency under 980 nm excitation, the RB emission band peaking at $\sim$590 nm in (B) is magnified by 50 times. (C) The integrated intensity of the RB emission band in (B), (D) ET efficiency of the six UCNP-RB samples. The emission spectra were normalized at 650 nm for each sample before calculation.

Figure 2. Bright-field SEM images and size distribution of NaYF$_4$:Yb$^{3+}$, Er$^{3+}$ core (A) and NaYF$_4$:Yb$^{3+}$, Er$^{3+}$/NaYF$_4$ core–shell UCNPs (B–F) corresponds to cs1 to cs5, respectively. The size distribution was given by averaging the diameter ± standard deviation (more than 300 nanoparticles were counted).

Figure 3. Emission spectra of UCNPs (A) and UCNP-RB samples (B) under 980 nm excitation. The RB emission band peaking at $\sim$590 nm in (B) is magnified by 50 times. (C) The integrated intensity of the RB emission band in (B), (D) ET efficiency of the six UCNP-RB samples. The emission spectra were normalized at 650 nm for each sample before calculation.
relaxation processes, and the emission kinetics is always faster in UCNP-RB samples than that in the corresponding UCNPs, whereas for 650 nm emission, the temporal behavior of UCNPs and corresponding UCNP-RB are almost the same as shown in Figure S2 and Table S2 (SI). It confirms the occurrence of dynamic ET in the green spectral region from UCNPs to RB molecules.

The amplitude ratio $A_2/A_3$ of the slow to the fast components of 540 nm emission in Table 1 gets larger with an increase of the shell thickness, indicating that more emitting centers are out of the dynamic ET interaction distance. It is consistent with the distance increase between the doped area of UCNPs and RB molecules. Assuming that the emitting centers are distributed evenly in the core of UCNPs, the two areas can be calculated from $A_2/A_3$, and the Förster distance was found to be shell-thickness-dependent (see eq S1 and the relevant analysis in the SI). For UCNPs with the thickest shell, the upconversion luminescence quantum efficiency is enhanced by ~36-fold, and accordingly, the Förster distance is about 1.8 times that of the bare core. Therefore, with this NaYF$_4$ shell on the surface of NaYF$_4$:Yb$^{3+}$, Er$^{3+}$ UCNPs, the donor–acceptor distance is increased, disabling dynamic ET; in the meantime, however, the dynamic ET interaction distance is increased due to the enhancement of upconversion luminescence, which is beneficial for dynamic ET performance. Therefore, an optimal shell thickness exists for dynamic ET as well.

Table 1. Fitted Time Constants for UCNP-RB Samples and the Amplitude Ratio for 540 ($A_2/A_3$) and 590 nm ($A_2'/A_3'$)

<table>
<thead>
<tr>
<th></th>
<th>core</th>
<th>cs1</th>
<th>cs2</th>
<th>cs3</th>
<th>cs4</th>
<th>cs5</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\tau_{rise, RB}/\mu$s</td>
<td>4.50</td>
<td>6.62</td>
<td>9.31</td>
<td>13.88</td>
<td>14.16</td>
<td>16.66</td>
</tr>
<tr>
<td>$\tau_2/\mu$s</td>
<td>113.71</td>
<td>136.50</td>
<td>187.08</td>
<td>255.59</td>
<td>276.50</td>
<td>327.39</td>
</tr>
<tr>
<td>$\tau_3/\mu$s</td>
<td>50.29</td>
<td>69.84</td>
<td>91.97</td>
<td>113.32</td>
<td>122.82</td>
<td>130.54</td>
</tr>
<tr>
<td>$A_2/A_3$</td>
<td>0.61</td>
<td>1.05</td>
<td>1.40</td>
<td>1.90</td>
<td>3.03</td>
<td>3.21</td>
</tr>
<tr>
<td>$A_2'/A_3'$</td>
<td>0.06</td>
<td>0.11</td>
<td>0.31</td>
<td>0.46</td>
<td>1.31</td>
<td>1.56</td>
</tr>
</tbody>
</table>

Table 2. Shell Thicknesses of the Six UCNPs and the Dynamic and Static ET Components for RB Excitation

<table>
<thead>
<tr>
<th></th>
<th>core</th>
<th>cs1</th>
<th>cs2</th>
<th>cs3</th>
<th>cs4</th>
<th>cs5</th>
</tr>
</thead>
<tbody>
<tr>
<td>shell thickness/nm</td>
<td>0</td>
<td>0.9</td>
<td>2.7</td>
<td>4.2</td>
<td>6.4</td>
<td>8.9</td>
</tr>
<tr>
<td>$D/S$</td>
<td>10.02</td>
<td>8.30</td>
<td>3.60</td>
<td>3.10</td>
<td>1.32</td>
<td>1.05</td>
</tr>
<tr>
<td>$I_{dynamic}/I_{total}$</td>
<td>0.80</td>
<td>0.71</td>
<td>0.45</td>
<td>0.32</td>
<td>0.10</td>
<td>0.07</td>
</tr>
<tr>
<td>$I_{static}/I_0$</td>
<td>0.80</td>
<td>1.42</td>
<td>3.15</td>
<td>3.84</td>
<td>1.00</td>
<td>0.63</td>
</tr>
<tr>
<td>$I_{dynamic}/I_0$</td>
<td>0.20</td>
<td>0.58</td>
<td>3.85</td>
<td>8.16</td>
<td>9.00</td>
<td>8.37</td>
</tr>
</tbody>
</table>

$I_0$ denotes the total emission of RB from the core-RB sample.

Figure 4. Time behavior of upconversion luminescence of UCNPs (red curves) and UCNP-RB (blue curves) at 540 nm and RB emission of UCNP-RB at 590 nm (magenta curves) under 980 nm excitation. The raw data are presented with black scatter, and the fitted results are shown with colored solid curves.
efficient dynamic or static ET occurs in the core–shell sample with more or less the same shell thickness where the most efficient ET (dynamic plus static ET) occurs.

In conclusion, aiming at optimizing the biofunctional upconversion nanoplatorms, the shell role in the interplay between static and dynamic ET mechanisms has been elucidated for the first time for upconversion nanosystems. The ET process was modeled and experimentally validated on a typical biofunctional upconversion nanoplatorm. From the obtained relation, it was determined that although the contribution of the dynamic part in ET decreases when the shell gets thicker, it reaches a maximum with a shell thickness of ~4 nm. On the contrary, the contribution of static ET increases when the shell gets thicker and reaches a maximum with a similar shell thickness. In a word, optimization of static ET, as well as dynamic ET, in biofunctional nanoplatorms can be achieved by coating a proper shell. This work is significant in optimization of upconversion nanoplatorms applied in, but not limited to, bioassays and photodynamic/thermal therapies.

**ASSOCIATED CONTENT**

**Experimental details.** Data analysis. XRD patterns of UCNPs. Time behavior of upconversion luminescence at 650 nm. The fitting results. The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcllett.5b00999.

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**Notes**  
The authors declare no competing financial interest.

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