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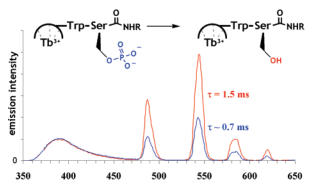
Phosphorylation State-Responsive Lanthanide Peptide Conjugates: A Luminescence Switch Based on Reversible Complex Reorganization

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ABSTRACT



A luminogenic probe for peptide dephosphorylation has been developed. It consists of a serine-/tyrosine-containing peptide modified on the N-terminus with a tryptophan residue and a DTPA chelate capable of binding Tb³⁺. We propose a mechanistic model for the luminescence enhancement based on the interconversion of monomeric and dimeric lanthanide species, which is affected by the phosphorylation state of the serine or tyrosine residue. The optical switch reports effectively on phosphatase-catalyzed dephosphorylation in vitro.

Probe molecules that relay information about chemical transformations via an optical response enable monitoring and imaging of these processes in complex systems, such as live cells.¹ A significant challenge in probing biological systems is the presence of native chromophores that emit UV/visible light on the nanosecond time scale ("cellular autofluorescence"). This is typically overcome by using fluorescent probes that emit at longer wavelengths than the native chromophores.¹ However, an additional degree of resolution can be achieved by transposing the signal output to the millisecond time domain using sensitized lanthanide luminescence.² In this context, lanthanide luminescence has

been used for studying the structure³ and metal-binding affinity of proteins⁴ as well as for sensing biologically relevant analytes⁵ and redox poise.⁶

Phosphorylation of proteins and peptides stands out among posttranslational modifications as the predominant mechanism of cellular signaling.⁷ Consequently, these processes have been targets of optical probing strategies. Although significant advances have recently been reported,⁸ robust

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optical switches for phosphorylated/dephosphorylated peptide pairs have been difficult to design owing to their relatively subtle structural differences.⁹

As part of a broad program aimed at sensing chemical/enzymatic transformations via optical probes, 10 we became interested in developing an optical switch based on the phosphorylated state of a peptide that would employ lanthanide luminescence and that would potentially be able to report on phosphorylation/dephosphorylation processes. We synthesized ligands 1 and p1, which contain a diethylenetriaminepentaacetic acid (DTPA) ligand, a tryptophan residue capable of sensitizing Tb^{3+} luminescence, 4b and serine and phosphoserine residues, respectively (Figure 1a). We inves-

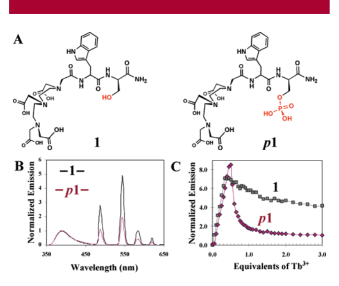


Figure 1. (a) Structures of **1** and $p\mathbf{1}$. (b) Steady-state luminescence spectra of **1** and $p\mathbf{1}$ in the presence of 1 equiv of TbCl₃. (c) Titrations of **1** (gray squares) and $p\mathbf{1}$ (purple diamonds) with TbCl₃ (emission monitored at 545 nm). Spectra are normalized to emission at 390 nm, which remains constant after \sim 1 equiv.

tigated their optical properties and found that in the presence of $\mathrm{Tb^{3+}}$ there was a significant difference in luminescence intensity (Figure 1b). We were surprised to see that p1 was less intense than 1 because the initial design was based on the premise that the coordination of the phosphoserine of p1 to $\mathrm{Tb^{3+}}$ should provide enhanced luminescence. We further investigated this behavior by titrating 1 and p1 with

Tb³⁺ (Figure 1c). Interestingly, these experiments showed that for both **1** and *p***1** luminescence maximized after the addition of \sim 0.5 equiv of Tb³⁺, suggesting the formation of a ternary (or higher order) adduct rather than the expected 1:1 Tb/DTPA—peptide complex.¹¹ The decrease in luminescence intensity after this maximum was more pronounced for *p***1** than for **1**, such that in the presence of 1–2 equiv of Tb³⁺ the two compounds were easily differentiated.

Time-resolved emission experiments were carried out on selected titration points (Figure 2). Luminescence decay

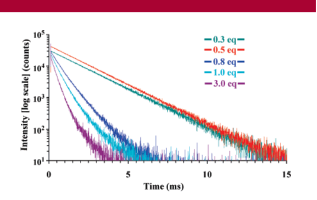


Figure 2. Luminescence decay curves for p1 in H₂O (100 μ M) with 0.3 (teal), 0.5 (red), 0.8 (blue), 1.0 (cyan), and 3.0 (purple) equiv of TbCl₃.

curves of *p*1 in the presence of 0.3 or 0.5 equiv of Tb³⁺ were monoexponential, indicating the presence of a single luminescent species with a lifetime of about 1.7 ms. At higher Tb³⁺ concentrations (0.8, 1.0, 3.0 equiv), the luminescence is markedly shorter lived, and the decay curves are no longer monoexponential, indicating the presence of multiple luminescent species. By contrast, the luminescence decay curves of 1 were observed to be monoexponential at all Tb³⁺ concentrations, although the lifetimes decreased with increasing Tb³⁺ concentration.¹⁶

Guided by both the steady-state and the time-resolved emission measurements, we propose the following model for the observed luminescence switch 1/p1. Each peptide probe (1 and p1) exists in solution containing Tb^{3+} (≥ 1 equiv) as an equilibrium of two species, the expected monomeric and the dimeric complex (Figure 3). The dimeric species (for both 1 and p1) is more luminescent than the corresponding monomer because it is likely that fewer quenching water molecules are bound to the metal center. The position of the equilibrium between these two species is affected by the phosphate group. Specifically, in the presence of the phosphate moiety, the equilibrium is shifted in favor of the monomeric p1/Tb complex, presumably due to intramo-

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⁽¹²⁾ Although higher-order adducts may be present in solution, the simplest ligand/metal stoichiometry that is consistent with the titration data is 2:1. Mass spectrometry in various ionization modes was unsuccessful in confirming the identity of the oligomeric species, although the monomeric p1 was readily observed; see Supporting Information for additional details.

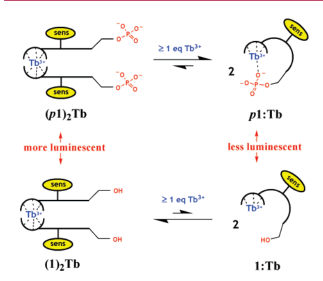


Figure 3. Proposed model for the observed luminescence switch 1/p1 based on reversible complex reorganization. The presence of phosphate shifts the equilibrium toward the less-luminescent monomeric p1:Tb species, presumably because of intramolecular interaction between the phosphate and the metal.

lecular interaction between the phosphate and the metal.¹³ It should be noted that it is likely that this behavior is the observed sum of a large number of interactions resulting from a variety of microenvironments fostered by the presence of the phosphate group. In contrast, in the absence of the phosphate, the equilibrium favors the dimeric complex.

Thus, the luminescence increase produced by dephosphorylation is a result of converting one equilibrium, which favors the monomer, to another equilibrium, which favors the more luminescent dimer. Although interconversion of mono- and bis-ligated lanthanide species is known to occur with tripodal aminomethylene tris(aryl) ligands, ¹⁴ there are no reports of such behavior with polyaminocarboxylates such as DTPA. ¹⁵

Further support for the dimerization of these peptide probes was provided by monitoring the Tb³⁺ titration of *p*1 by ³¹P NMR.¹⁶ The size of the peak at 0.5 ppm corresponding to the phosphoserine residue decreases linearly as Tb³⁺ is added and is completely consumed after the addition of 0.5–0.6 equiv. The disappearance of the peak is consistent with the incorporation of the phosphopeptide into a Tb³⁺ complex, which is known to cause extreme peak broadening.^{4a} The same trend was observed for a phosphotyrosine analogue of *p*1. Unfortunately, no new phosphorus signal was observed after the disappearance of the free ligand peak (>0.5 equiv

of Tb³⁺), which prevented us from obtaining direct evidence for the proposed phosphoserine coordination to Tb³⁺ in the monomeric species.¹⁷

According to this mechanistic hypothesis, analogues of peptides 1/p1 containing other substituents capable of chelating the oxophilic, cationic metal center should also show a similar luminescence switching. This prediction was confirmed by the following examples (Figure 4). The tyrosine/

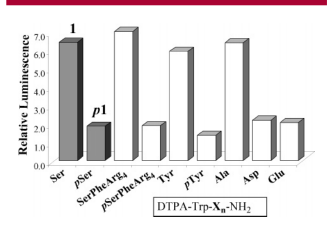


Figure 4. Tb/Trp emission ratios (545/390 nm) for analogous DTPA peptide conjugates (50 μ M in 100 μ M TbCl₃, 10 mM HEPES, 100 mM NaCl, pH 7.4, 25 °C). Peptides with anionic residues in the X_n position show decreased luminescence, supporting the intramolecular interaction rationale.

phosphotyrosine peptide pair exhibits luminescence properties very similar to those for 1/p1. Also, peptides containing a carboxylate on the residue adjacent to the DTPA-Trp fragment showed less intense Tb³⁺ luminescence than those containing neutral functionalities at this position; for example, DTPA-TrpAspNH₂ and DTPA-TrpAlaNH₂ constitute an optical switch as well. Furthermore, the phospho-/dephosphopeptide pair DTPA-Trp(p)SerPheArg₄NH₂ exhibited switching behavior nearly identical to the parent pair 1/p1, despite the presence of four cationic arginine residues, suggesting that interaction between the peptide chains is not a contributing factor (Figure 4).

To assess the ability of this optical switch to serve as a luminogenic probe of an enzymatic transformation, p1 was subjected to the action of alkaline phosphatase, a promiscuous enzyme capable of hydrolyzing phosphopeptides. Indeed, at 37 °C in pH 7.4 HEPES buffer, the expected luminescence increase was observed and confirmed to correlate with conversion of p1 to 1 by mass spectrometry. ¹⁶ As a result, these probes allow for continuous (real-time) monitoring of peptide dephosphorylation in vitro. According to the mechanistic rationale discussed above, phosphatase converts one equilibrium to another, increasing the relative concentration of the more luminescent dimer complex (Figure 5).

In summary, we have presented a new format for discriminating between phosphorylated and dephosphorylated

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⁽¹³⁾ Titration of p1 at various ionic strengths showed that the sharp maximum could be "salted out", which is consistent with the switching phenomenon involving electrostatics.

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⁽¹⁷⁾ It is important to note that proximity to ${\rm Tb}^{3+}$, not only coordination to it, can cause extreme broadening of NMR signals. For more details, see Supporting Information.

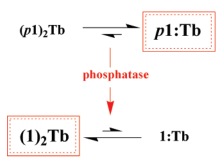


Figure 5. Schematic representation of the phosphatase-mediated conversion of p1 to 1.

peptides based on a significant change in luminescence properties. A mechanistic model for this behavior invokes the interconversion of mono- and bis-ligated lanthanide species, where the presence of a phosphate group stabilizes the monomeric complex. Although this luminescence switch reports adequately on a dynamic enzymatic process in vitro, its application in vivo may be limited because of the changes in complexation stoichiometry.

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Supporting Information Available: Experimental procedures, steady-state and time-resolved luminescence studies, and MS characterization. This material is available free of charge via the Internet at http://pubs.acs.org.

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