The origins of binding specificity of a lanthanide ion binding peptide

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Lanthanide ions (Ln3+) show similar physicochemical properties in aqueous solutions, wherein they exist as +3 cations and exhibit ionic radii differences of less than 0.26 Å. A flexible linear peptide lanthanide binding tag (LBT), which recognizes a series of 15 Ln3+, shows an interesting characteristic in binding specificity, i.e., binding affinity biphasically changes with an increase in the atomic number, and shows a greater than 60-fold affinity difference between the highest and lowest values. Herein, by combining experimental and computational investigations, we gain deep insight into the reaction mechanism underlying the specificity of LBT3, an LBT mutant, toward Ln3+. Our results clearly show that LBT3-Ln3+ binding can be divided into three, and the large affinity difference is based on the ability of Ln3+ in a complex to be directly coordinated with a water molecule. When the LBT3 recognizes a Ln3+ with a larger ionic radius (La3+ to Sm3+), a water molecule can interact with Ln3+ directly. This extra water molecule infiltrates the complex and induces dissociation of the Asn5 sidechain (one of the coordinates) from Ln3+, resulting in a destabilizing complex and low affinity. Conversely, with recognition of smaller Ln3+ (Sm3+ to Yb3+), the LBT3 completely surrounds the ions and constructs a stable high affinity complex. Moreover, when the LBT3 recognizes the smallest Ln3+, namely Lu3+, although it completely surrounds Lu3+, an entropically unfavorable phenomenon specifically occurs, resulting in lower affinity than that of Yb3+. Our findings will be useful for the design of molecules that enable the distinction of sub-angstrom size differences.

Lanthanide elements (Ln) consist of 15 elements with similar physicochemical properties; they exist as +3 cations (Ln3+) in solution and exhibit ionic radii differences of less than 0.26 Å1. Because of their unique magnetic and optical properties, Ln-element-based compounds are applied in various advanced materials, such as rechargeable batteries, lamp phosphors, and permanent magnets2–5. For sustainable usage and advancement of Ln elements, the construction of more efficient recovery techniques and recycling methods is required4–6. Therefore, many researchers have attempted to find molecules with recognition ability toward each Ln element6–8. For example, some chelating agents [e.g., ethylenediaminetetraacetic acid (EDTA) and nitrilotriacetic acid (NTA)] have been demonstrated to bind with Ln3+, and their affinity gradually increases as the ionic radius of Ln3+ decreases (Supplemental Fig. S1a,b)9,10. This gradual increase in affinity is simply explained by the trend of increasing Ln3+ acidity with decreasing ionic radius. However, most of these molecules lack the recognition specificity for clearly identifying individual Ln3+ elements.

A linear peptide consisting of 17 amino acids, named lanthanide binding tag (LBT), shows an interesting recognition pattern for Ln3+.1 This peptide originates from a calcium-binding protein, calmodulin, and has been engineered to recognize Ln3+ by combinatorial screening12,13. Because of the short peptide sequence and high binding affinity, LBT has been incorporated into recombinant proteins and used as an effective tool for structural analysis14–17. In particular, the binding affinity of LBTv, one of the LBT variants, along the Ln3+ series does not correlate simply to the decrease in the ionic radii (Supplemental Fig. S1c); a biphasic change occurs with Eu3+ as the branching point and shows drastically different binding affinities toward different Ln3+, exhibiting a > 60-fold lower affinity for La3+ than for Tb3+, with KD values of 3500 nM and 57 nM, respectively11. These binding properties indicate that a different mechanism underlies the recognition specificity of LBT compared to that of other chelating agents such as EDTA and NTA; however, the origin of this recognition specificity has not yet been investigated. The elucidation of the recognition mechanism of LBT toward Ln3+ would greatly contribute to the development of specific agents for Ln3+ recovery as well as for the accurate artificial design of functional peptides.

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Herein, to understand the mechanism of LBT specificity, we evaluated the thermodynamic parameters of binding between LBT3, an LBT derivative exhibiting the highest binding affinity to Ln$^{3+}$, and Ln$^{3+}$. We further elucidated the structures of LBT3-Ln complexes in solution. In addition, the structural fluctuations and the interactions between the complexes and the surrounding water molecules were analyzed using molecular dynamics (MD) simulations. Our results showed that the structures of the LBT3-Ln complexes for different Ln$^{3+}$ are similar. However, the thermodynamic parameters ($\Delta H$ and $\Delta S$) changed intricately across the series of Ln$^{3+}$, which can be divided into three categories, namely, La$^{3+}$ to Sm$^{3+}$, Sm$^{3+}$ to Yb$^{3+}$, and Lu$^{3+}$. In addition, we clarified that the large affinity difference is attributed to a water molecule, which directly coordinates with Ln$^{3+}$ in an LBT3-Ln complex.

**Results**

**Thermodynamic parameters of LBT3 binding with Ln$^{3+}$**. Isothermal titration calorimetry (ITC) experiments were performed to evaluate the thermodynamic parameters of the binding of LBT3 with a series of Ln$^{3+}$ (Fig. 1, Supplemental Table S1). All experiments were performed at pH 6.0, as some Ln$^{3+}$ easily form hydroxide species and precipitate above pH 7.0.$^9$ All the reactions showed endothermic behavior; the change in free energy ($\Delta G$) decreased with increasing Ln$^{3+}$ atomic number, from La$^{3+}$ ($\Delta G = -6.9$ kcal/mol) to Tb$^{3+}$ ($\Delta G = -9.1$ kcal/mol). The $\Delta G$ then remained almost constant from Tb$^{3+}$ to Yb$^{3+}$, and then increased slightly for Lu$^{3+}$. Due to the differences in measurement conditions, especially pH and temperature, the current affinity values show an approximately tenfold difference compared to those in previous reports using the same LBT$^{33,34}$. However, the relative affinities for each Ln$^{3+}$ were the same as those of LBTv (Supplemental Fig. S1e)$^{11}$.

The changes in the enthalpy ($\Delta H$) and entropy ($\Delta S$) of binding with each Ln$^{3+}$ followed a different pattern than that for $\Delta G$; both parameters decreased from La$^{3+}$ to Sm$^{3+}$, increased from Sm$^{3+}$ to Yb$^{3+}$, and finally decreased again for Lu$^{3+}$. These results indicate that the difference in affinity is not solely dependent on the acidity of the Ln$^{3+}$, but could be influenced by multiple factors, such as the ion size, hydration number, and acidity of the Ln$^{3+}$.

Moreover, we surmised that these differences might be responsible for the structural differences among the LBT3-Ln complexes.

**Differences between the $^1$H NMR spectra of the LBT3-La and LBT3-Lu complexes.** To compare the structures of a strongly bound and weakly bound complex, La$^{3+}$ and Lu$^{3+}$ were chosen for nuclear magnetic resonance (NMR) experiments. LBT3 exhibits a greater than 40-fold difference in affinity toward La$^{3+}$ and Lu$^{3+}$ under our conditions (Supplemental Table S1)$^{11}$, which might reflect a difference in the LBT3-Ln structures. Moreover, these two ions are diamagnetic, which enables a straightforward comparison of their NMR spectra. The other Ln$^{3+}$ in the series are paramagnetic, which leads to drastic changes in chemical shifts and peak shapes.$^{18}$ As shown in Fig. 2, there are obvious differences between the $^1$H NMR spectra of the LBT3-La and LBT3-Lu complexes; LBT3-Lu displays sharp peaks, while LBT3-La exhibits broadened peaks; large chemical shift differences were observed in the amide protons of N5 (N5-HN) and G6 (G6-HN), of 0.5 ppm and 0.7 ppm, respectively (Supplemental Fig. S2). Upon titration, we observed a clear difference between the exchange rates of the binding of LBT3 with La$^{3+}$ and Lu$^{3+}$; a slow exchange was observed for Lu$^{3+}$, and an intermediate exchange was observed for La$^{3+}$ (Supplemental Fig. S3). Titration of Lu$^{3+}$ into the LBT3 solution resulted in the immediate formation of new peaks related to complexation, while the peak intensity of free LBT3 decreased as the Lu$^{3+}$ concentration increased. The free LBT3 peak disappeared upon the addition of a two-fold Lu$^{3+}$ concentration. Conversely, titration with La$^{3+}$ initially induced a decrease in the intensity of the free LBT3 peaks. An unknown peak was observed at LBT3:La = 1:1, and peaks associated with complex formation resolved at LBT3:La = 1:2. Variable temperature $^1$H NMR measurements also showed that the LBT3-La spectrum sharpened at higher temperatures, indicating an intermediate to fast exchange, while the LBT3-Lu peaks became broad, indicating a slow to intermediate exchange (Supplemental Fig. S4).

**Structures of LBT3-Ln complexes.** To elucidate the structures of LBT3-Ln complexes in aqueous solution, gel-filtration chromatography was first performed to determine the self-assembled state at pH 6.0. As a result, free LBT3 showed a clear single peak at the three concentrations tested, and no differences were observed in the retention volume (Supplemental Fig. S5a). Moreover, the complexes LBT3-La and LBT3-Lu both showed clear single peaks and larger retention volumes, compared to free LBT3 (Supplemental Fig. S5b). These results...
assignments are based on standard homonuclear two-dimensional NMR methodology. In this case, the distance between N5-Oδ1 and La3+ rapidly increased from ~0.2 nm to >0.4 nm for the 315 K and 345 K calculations (Fig. 4, Supplemental Fig. S9). All the other coordinated residues consistently chelated Ln3+ throughout the calculations in both complexes (Supplemental Fig. S9c,d). Although the N5-O61 dissociation is only observed in 3 out of 10 conditions examined here (possibly due to the slightly shorter calculation time), these results, at least, indicate that the binding between N5-O61 and Ln3+ was considerably weaker than that of the other coordinating residues, especially in the case of La3+. This is reasonable because this dipole-ion interaction is weaker than the ion-ion interactions. Although the continuously bound site W9O also coordinated Ln3+ via dipole-ion interactions, W9O is tightly structurally restrained compared to the other residues.

The free energy landscape as a function of the distance between N5-O61 and Ln3+ was calculated using the accelerated weight histogram (AWH) method for 200 ns each at 300 K. To distinguish the association/dissociation effect of N5-O61 from the effect of a large overall structural change in LBT3, 0.5 kcal/mol/Å² harmonic constraints were applied to the backbones during the free energy calculations. Remarkable differences were observed between the free energy landscapes of the 6LBT3-La and 6LBT3-Lu systems. In the 6LBT3-Lu system, the associated state is 7.1 kcal/mol more stable than the dissociated state, whereas in 6LBT3-La, the associated state is 7.1 kcal/mol more stable than the dissociated state, whereas in 6LBT3-La, the associated state is 7.1 kcal/mol more stable than the dissociated state.
state was only 4.6 kcal/mol more stable than the dissociated state (Fig. 4c). Although artificial constraints were applied in these simulations, these calculations also indicated that the dissociation of N5-Oδ1 occurs more easily in LBT3-La than in LBT3-Lu.

Monodentate or bidentate chelation between carboxylate oxygen and Ln³⁺. In addition to N5-Oδ1, D3 and D7 carboxylate groups showed a difference in recognition between La³⁺ and Lu³⁺ (Fig. 5a, b, Supplemental Fig. S9c,d). Although all carboxylate groups remained bound to Ln³⁺ throughout the 100 ns calculations under all temperature conditions, D3 and D7 carboxylate oxygens frequently showed bidentate chelation in the 6LBT3-La system, whereas monodentate chelation of these coordinates was observed for 6LBT3-Lu. In the case of monodentate binding, the carboxylate oxygens bind with Ln³⁺ and a nearby water molecule. It is considered that this observation relates to ion size. To confirm this hypothesis, additional MD simulations were performed; the La³⁺ in the 6LBT3-La complex was replaced with Ln³⁺. The results showed that the distance between D3-Oγ1 and Ln³⁺ is shortened as the ionic radii decrease. In contrast, the distance between D3-Oγ2 and Ln³⁺ is shortened from La³⁺ to Sm³⁺, increased from Sm³⁺ to Tb³⁺, and finally becomes insignificant from Tb³⁺ to Lu³⁺ (Fig. 5c). The D7 carboxylate also showed that D7-Oγ1 is simply nearing Ln³⁺ as the ionic radii decrease.

Figure 3. Stereo view of the complex structures of 6LBT3-La and 6LBT3-Lu. (a) 6LBT3-La (red) and 6LBT3-Lu (blue) are superimposed. The sphere represents Ln³⁺. (b) 180° rotated view of (a). 20 structures of each 6LBT3-Ln complex are superimposed. The sidechain hydrogens are omitted for clarity.
(Fig. 5d), while the average distance between D7-Oγ2 and Ln3+ is shortened from La3+ to Tb3+, and becomes insignificant from Tb3+ to Lu3+. In addition, the large error bars observed in Eu3+ to Lu3+ indicate that D7-Oγ2 easily dissociates from Ln3+ as the ionic radii decrease. These observations indicate that the ion size affects the orientation of the coordinated carboxylate oxygens.

**Direct water molecule coordination to Ln3+.** To evaluate the origin of the N5-Oδ1 coordination/dissociation, we closely analyzed the local structural changes around Ln3+. The coordination patterns of the 6LBT3-La and 6LBT3-Lu systems differed in their interaction with water molecules. Specifically, the 6LBT3-La systems exhibited four coordination patterns: (i) La3+ bound only to six LBT3 residues (Fig. 6a); (ii) La3+ bound directly to a water molecule in addition to six LBT3 residues (Fig. 6b); (iii) La3+ coordinated to five LBT3 residues (excluding N5-Oδ1) and a water molecule (Fig. 6c); and (iv) La3+ coordinated to five LBT3 residues (excluding N5-Oδ1) and two water molecules (Fig. 6d). During the 100 ns simulations at various temperatures, the (i) and...
(ii) states were frequently observed, and the coordination pattern changed as follows: (i) ⇄ (ii) ⇄ (iii) ⇄ (iv). Notably, the coordinated water molecule in (iii) was stabilized by hydrogen bonding with N5-Oδ1 (Fig. 6c). This water molecule invaded the packed complex structure, which led to the coordination of an additional water molecule to La³⁺, resulting in the (iv) state. Moreover, the α helical c-terminus observed in the (i) and (ii) state formed a stretched structure in the (iii) and (iv) state. The 6LBT3-Lu system, on the other hand, showed three types of coordination: Forms analogous to (i) and (iii) above, as well as a unique form in which (v) Lu³⁺ was coordinated with five residues (excluding N5-Oδ1), without a coordinated water molecule (Fig. 6e). The (ii) and (iv) forms were not observed for 6LBT3-Lu. The (i) form was highly stable; coordination changes were only observed above 345 K and followed the path (i) ⇄ (v) ⇄ (iii). The radial distribution functions (RDFs) between water and Ln³⁺ also indicated that the mechanistic differences between the two complexes originated at the point

Figure 6. Coordination patterns of LBT3-Ln observed in MD simulations. (a) Ln³⁺ bound to six LBT3 residues only. (b) Ln³⁺ bound to six LBT3 residues and a water molecule. (c) Ln³⁺ bound to five LBT3 residues (excluding N5-Oδ1) and a water molecule. (d) Ln³⁺ bound to five LBT3 residues and two water molecules. (e) Ln³⁺ bound to five LBT3 residues only. Ln³⁺ are indicated by dark green spheres and water molecules are indicated by red and white spheres (oxygen and hydrogen, respectively). Peptide backbones are shown by solid black lines. Red lines indicate oxygen. All hydrogen molecules of the peptide were omitted for clarity. Yellow arrows indicate the water molecule that is bound to Ln³⁺, while the blue arrow indicates the water molecule that is bound to D7-Oy2.
at which water coordination occurred. Direct coordination of water was observed under all conditions tested for 6LBT3-La, whereas it was only observed at 345 K for 6LBT3-Lu (Supplemental Fig. S10a,b). In addition, to check the initial structural effects, La3+ and Lu3+ were exchanged with each other, and MD simulations were performed at 300 K and 315 K. This showed that direct coordination of water was only observed in the 6LBT3-La complex, even when La3+ and Lu3+ were exchanged in the initial structures of the simulation (Supplemental Fig. S10c,d). These observations suggest that the direct water coordination strongly depends on the properties of the individual Ln3+, and should be related to the difference in size between the ions, rather than being an artifact derived from the initial structural differences.

NMR spectroscopy also supported the simulated results described above. Proton exchange was observed for N5-HN and N6-HN in the LBT3-La complex at 10 °C and 25 °C (Fig. 7a). The LBT3-Sm complex also showed proton exchange for N5-HN and N6-HN at 25 °C (Fig. 7b), but no amide proton exchange was observed for the LBT3-Lu complex (Fig. 7c). Proton exchange reflects the degree of solvent accessibility and mobility of each site. Considering the structures, water molecules are thought to exist inside the LBT3-La complex. These results also agreed well with the 1H NMR spectral differences between LBT3-La and LBT3-Lu; N5-HN of LBT3-La exhibited a comparatively large shift toward the higher magnetic field, 0.5 ppm, compared with LBT3-Lu (Supplemental Fig. S2). To confirm that the largest chemical shift observed for G8-HN is also induced by water molecule interaction, the RDFs between water and G8-HN were analyzed. A clear difference was observed, namely that a water molecule locates closer to the G8-NH of LBT3-La than to that of LBT3-Lu (Supplemental Fig. S11a,b). The detailed structural analysis showed that the difference originates from the orientation of the D3 carboxylate oxygen. When the D3 carboxylate exhibits bidentate chelation, a water molecule draws close to G8-HN (Supplemental Fig. S11c). In the case of monodentate chelation, the freed carboxylate oxygen (D3-Oγ2) traps a water molecule, preventing the water molecule from drawing close to G8-HN (Supplemental Fig. S11d). These experimental and computational results demonstrate that there is a notable difference between the interaction of water molecules with LBT3-La and LBT3-Lu. In particular, the water infiltration easily occurs in the case of the La3+ complexation, whereas this is a rare event in the Lu3+ recognition.
Effect of N5-Oδ1 dissociation to the complex structure. To elucidate the effect of N5-Oδ1 dissociation from Ln³⁺, we performed another MD simulation using artificial complex structures, where which Ln³⁺ interacts with the five sites (excluding N5-Oδ1) of LBT3. The initial structures were constructed using the program CYANA based on the NOESY spectra; the five residues were set as binding sites. The obtained structures, which were named 5LBT3-La and 5LBT3-Lu, showed high similarity with a backbone rmsd value of 0.456 Å (Supplemental Fig. S12). Using these structures, we performed MD simulations under the same conditions as for 6LBT3-Ln. Although N5-Oδ1 did not bind with Ln³⁺ at any point during the calculation (Supplemental Fig. S13a,b), the other coordination sites maintained their binding with Ln³⁺, similarly to the 6LBT3-Ln system (Supplemental Fig. S13c,d). In the 5LBT3-La system, as expected, direct water coordination was also frequently observed (Supplemental Fig. 14a). Structure (iii), which includes direct coordination of one water molecule to La³⁺, was the most frequently observed throughout the calculation at all the tested temperatures. The (iii) ⇄ (iv) structural transition was occasionally observed. By contrast, in the 5LBT3-Lu system, direct water coordination was only observed at higher temperatures (≥ 315 K) (Supplemental Fig. 14b). Only structure (v), which does not include direct coordination of any water molecule to Lu³⁺, was observed at low temperatures (≤ 300 K), and the coordination transition (v) ⇄ (iii) was observed at higher temperatures (≥ 315 K). Despite the artificial constraints in the initial structures of 5LBT3-Ln, these results indicate that when N5-Oδ1 is in the dissociation state, La³⁺ is exposed in the solvent, whereas Lu³⁺ is highly covered in LBT3.

Discussion

The overall binding pattern between LBT3 and Ln³⁺ is as follows. In aqueous solution, free LBT3 and Ln³⁺ are hydrated or coordinated, respectively, by water molecules that are released to the bulk solvent upon complex formation. LBT3 surrounds the Ln³⁺ by using its six residues and makes a partially helical compact structure. According to the experimental and computational results, the structural factor underlying the large difference in binding affinity of LBT3 with each Ln³⁺ can be explained as follows. From Sm³⁺ to La³⁺, as the coordination sphere enlarges, water molecules can directly interact with Ln³⁺ with ease. Because the coordinated water molecule is located next to N5-Oδ1, which is the most weakly coordinated among the ligands, it interposes itself between N5-Oδ1 and Ln³⁺. In addition, this water molecule invasion allows another water molecule to coordinate with Ln³⁺ directly; these water molecules interfere with the rebinding of N5-Oδ1 to Ln³⁺. The dissociated N5-Oδ1 results in the structural fluctuations of the N5 sidechain, and this local fluctuation spread may result in the structural disruption of the LBT3-Ln complex. In other words, the complexes with water molecules directly bound to Ln³⁺ are structurally flexible, resulting in a reduction in binding affinities. By contrast, from Sm³⁺ to Lu³⁺, the ion sizes are thought to be small enough to be completely covered by LBT3. Here, direct interaction with water is rare, resulting in stable complexation and high affinity.

The thermodynamic analysis indicates the magnitude of energies of various reactions during complexation. The water release event is enthalpically unfavorable, especially for such a large trivalent ion, and is not compensated by the enthalpically favorable reactions such as electrostatic bond formation between LBT3 and Ln³⁺, which results in a stable 8 or 9 coordination structure, and intramolecular hydrogen bond formation. Moreover, complex formation decreases the entropy of the peptide chain, which also disfavors binding. However, the release of water is very entropically favorable and represents the driving force of the reaction, ultimately overcoming any thermodynamically unfavorable effects. Although it is difficult to isolate various reactions and these energies, the structural features and thermodynamic parameters of each of the LBT3-Ln complexes allow us to interpret the recognition specificity as follows.

Considering the ΔH, ΔS (Supplemental Table S1) and structural features, the binding of LBT3 with Ln³⁺ could be divided into three: the first is La³⁺ ≤ Ln³⁺ ≤ Sm³⁺ (‘≤’ indicates the largeness of the atomic number), the second is Sm³⁺ ≤ Ln³⁺ ≤ Yb³⁺, and third is Lu³⁺ (Fig. 8). From La³⁺ to Sm³⁺ (top of Fig. 8), the complexes are less stable than the other classes. NMR and MD simulation results indicate that the LBT3 ligands are pulled tightly to Ln³⁺ directly; these water molecules interfere with the rebinding of N5-Oδ1 to Ln³⁺. The dissociated N5-Oδ1 results in the structural fluctuations of the N5 sidechain, and this local fluctuation spread may result in the structural disruption of the LBT3-Ln complex. In other words, the complexes with water molecules directly bound to Ln³⁺ are structurally flexible, resulting in a reduction in binding affinities. By contrast, from Sm³⁺ to Lu³⁺, the ion sizes are thought to be small enough to be completely covered by LBT3. Here, direct interaction with water is rare, resulting in stable complexation and high affinity.

As a result, the enthalpically disfavoring N5-Oδ1 dissociation becomes less, and the LBT3 ligands are pulled tightly to Ln³⁺ because of the increase in acidity. As a result, the enthalpically disfavoring N5-Oδ1 dissociation becomes less, and the intramolecular bond formation is induced, resulting in a decrease in ΔH. As a result, the enthalpically disfavoring N5-Oδ1 dissociation becomes less, and the intramolecular bond formation is induced, resulting in a decrease in ΔH. As a result, the enthalpically disfavoring N5-Oδ1 dissociation becomes less, and the intramolecular bond formation is induced, resulting in a decrease in ΔH. As a result, the enthalpically disfavoring N5-Oδ1 dissociation becomes less, and the intramolecular bond formation is induced, resulting in a decrease in ΔH. As a result, the enthalpically disfavoring N5-Oδ1 dissociation becomes less, and the intramolecular bond formation is induced, resulting in a decrease in ΔH.
strongly attracts the coordination atoms, although it also simultaneously induces closer coordination, which should result in repulsion. The intricately folded peptide structure might be distorted by the strong attraction. It was considered that the observed orientation change of the coordinated carboxylates of D3 and D7 results from the compensation of these steric hindrances.

Regarding Lu$^{3+}$ (bottom of Fig. 8), the LBT3-Lu complex is also occupied by the (i) form. Because of its small ion size, Lu$^{3+}$ is known to coordinate with notably fewer water molecules among the series of Ln$^{3+}$. Therefore, when LBT3 binds with Lu$^{3+}$, it needs less energy for the de-coordination of water molecules than Yb$^{3+}$. This also means that Lu$^{3+}$ releases fewer water molecules than Yb$^{3+}$. The decrease in $\Delta H (\Delta\Delta H_{\text{Lu-Yb}} = -0.4 \text{ kcal/mol})$ and $\Delta S (\Delta\Delta S_{\text{Lu-Yb}} = -0.63 \text{ kcal/mol})$ is predicted to reflect this character.

In conclusion, this is the first report that precisely explains how a flexible linear peptide LBT3 recognizes a Ln$^{3+}$ species. Our findings clearly indicate that water molecules play important roles, both in the reaction as a whole, and in the recognition specificity. Previously, the contributions of water molecules in protein-target interactions have been well studied. For example, in the potassium ion channel, which recognizes the difference between potassium and sodium ions, the protein structure controls the coordination/dissociation of water molecules around the ions. This is the key factor in distinguishing between these two ions, whose radii differ by less than 0.5 Å. Interestingly, a highly stabilized membrane protein and a flexible peptide use water molecules to distinguish sub-angstrom size differences. This finding could be important for the design of artificial molecules that considers solvent molecules as a part of the target molecule rather than solely as a solvent.

Experimental procedures

**Materials.** All lanthanide nitrates were purchased from Sigma-Aldrich (St. Louis, MO). MES-$d_{13}$ was purchased from Cambridge Isotope Laboratories Inc. (Tewksbury, MA).

**ITC analysis.** The thermodynamic parameters of binding between Ln$^{3+}$ and the peptides were analyzed at 10 °C by isothermal titration calorimetry (ITC, MicroCal VP-ITC, MicroCal Worcestershire, UK). The synthetic peptides and lanthanide nitrate (Ln(NO$_3$)$_3$) were each dissolved in 50 mM MES buffer containing 100 mM NaCl (pH 6.0). The experimental conditions were adjusted according to the manufacturer's instructions. Typically, LBT3 solution (15 μM) was placed in the calorimeter cell, and then Ln(NO$_3$)$_3$ (300 μM) was loaded into the syringe injector. The experiments included 20 injections, whereby an initial 2 μL injection was used to account for dilution of the syringe, and the remaining injections were 10 μL with a 300 s delay between each injection. The effect of Ln$^{3+}$ dilution in the cell was calculated by subtraction of titration data for a blank, which consisted of
titrating Ln3+ into a buffer solution. Each binding parameter, including stoichiometry (N), association constant (K), and binding enthalpy (∆H), was calculated using the ITC Origin Analysis Software version 7.0 (Malvern). For the samples that did not show a sigmoidal response owing to a low K value, the thermodynamic parameters were calculated by fixing the stoichiometry to 1.0 according to the manufacturer’s instructions.

**NMR spectroscopy.** Nuclear magnetic resonance (NMR) experiments were carried out using a Bruker Avance spectrometer equipped with a TCI cryoprobe. Standard 5 mm NMR tubes were used for the measurements. The samples were prepared in 30 mM MES-d13 buffer (H2O/D2O = 90:10, pH 6.0, Sigma-Aldrich). To equilibrate the binding reaction, all measurements were performed at least 10 min after mixing. All assignments were carried out using a combination of total correlation spectroscopy (TOCSY), nuclear Overhauser effect spectroscopy (NOESY), heteronuclear single-q quantum correlation spectroscopy (HSQC)27. In addition, CLEAN chemical exchange spectroscopy (CLEANEX-PM) was carried out to examine the solvent-accessible residues28. TOCSY was conducted with a mixing time of 80 ms, NOESY with a mixing time of 400 ms, and CLEAN-EX-PM with a mixing time of 200 ms. Titration experiments were performed by adding Ln3+ solutions to the peptide solutions. 1H and TOCSY spectra were measured at each titration step to trace the chemical shift changes. All NMR data were processed and analyzed with Topspin 3.1, NMRPipe29, and Sparky30.

**Structural calculation.** The peptide structures were calculated using the program CYANA 2.1 for the automatic assignment of the NOE peak lists. The upper limits of the distance restraints were calculated from the NOE cross-peak intensities using the calibration routine of CYANA. For calculation of the 6LBT3-Ln complex structures, the coordination oxygen atoms, the upper and lower limit distance restraints between Ln3+ to coordinated oxygen atoms were set according to the crystal structure of LBTv-Tb; D3-Oγ1, N5-Oδ1, D7-Oγ1, W90, E11-Oε1, E11-Oε2, E14-Oε1, and E14-Oε2 were set as coordination sites; the upper limit was set to 3.0 Å, and the lower limit was set to 2.1 Å. For calculation of the 5LBT3-Ln complex structures, D3-Oy1, D7-Oy1, W90, E11-Oε1, E11-Oε2, E14-Oε1, and E14-Oε2 were set as coordination sites. The structure with the lowest target function was used in subsequent MD simulations.

For the structural refinement, GROMACS software was used. Briefly, the 20 structures obtained by CYANA structural calculation were treated with the same procedure as subsequent MD simulation analyses until the production run. After 100 ps constant pressure equilibration, MD simulations with distance restraints were performed at 500 K for 10,000 steps. The structures were then minimized at the end of calculations. The structural representation was performed by MOLMOL 1.07 software31.

**Molecular Dynamics Simulations and Analysis.** All MD simulations were performed using GROMACS software32. Each LBT3-Ln complex was solvated in a square box of 2589 water molecules. All Glu and Asp residues were set as negatively charged forms, whereas the other residues were set to neutral. As a result, the net charge of the LBT3-Ln complex became -2, and Na+ counterions were added to neutralize the net charge. CHARMM36 force fields were employed for the peptide model33. TIP3P was used for water molecules, and the model of Migliorati et al.34 was used for the Ln3+. The long-range electrostatic interactions were treated with particle-mesh Ewald summation35, and a 1.2 nm cutoff was used for Lennard–Jones and Coulombic interactions. All bonds between hydrogen and heavy atoms were constrained using the LINCS algorithm36.

In the simulations, we initially performed 50,000 steps of minimizations with a restricted LBT3-Ln complex and then equilibrated the minimized systems following 100 ps constant pressure equilibrations. After equilibration, an additional 100 ps constant-volume equilibration and 100 ps constant pressure equilibration were performed without restrictions. The following production runs were then carried out for 100 ns. The temperature was kept constant using the stochastic velocity rescaling method37, and the pressure was kept constant at 1 atm using the Parrinello–Rahman method38. A 2 fs time step was employed in all simulations. Other simulation setups were followed according to the manufacturer’s instructions and the main text of our paper.

**Data availability**

The assigned 1H and 13C chemical shifts of LBT3-Ln have been deposited in the BMRB (36356, 36357) and structural coordinates have been deposited in the PDB (7CCN, 7CCO). All data are contained within the manuscript.

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Competing interests
The authors declare no competing interests.

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