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Observation of μ s time-scale protein dynamics in the presence of Ln^{3+} ions: application to the N-terminal domain of cardiac troponin C

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Abstract The microsecond time-scale motions in the N-terminal domain of cardiac troponin C (NcTnC) loaded with lanthanide ions have been investigated by means of a ¹H^N off-resonance spin-lock experiment. The observed relaxation dispersion effects strongly increase along the series of NcTnC samples containing La³⁺, Ce³⁺, and Pr³⁺ ions. This rise in dispersion effects is due to modulation of long-range pseudocontact shifts by μ s time-scale dynamics. Specifically, the motion in the coordination sphere of the lanthanide ion (i.e. in the NcTnC EF-hand motif) causes modulation of the paramagnetic susceptibility tensor which, in turn, causes modulation of pseudocontact shifts. It is also probable that opening/closing dynamics, previously identified in Ca²⁺-NcTnC, contributes to some of the observed dispersions. On the other hand, it is unlikely that monomer-dimer exchange in the solution of NcTnC is directly responsible for the dispersion effects. Finally, on-off exchange of the lanthanide ion does not seem to play any significant role. The amplification of dispersion effects by Ln³⁺ ions is a potentially useful tool for studies of µs-ms motions in proteins. This approach makes it possible to observe the dispersions even when the local

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Present Address: C. Eichmüller Sandoz GmbH, Biochemiestraße 10, A-6250 Kundl, Austria environment of the reporting spin does not change. This happens, for example, when the motion involves a 'rigid' structural unit such as individual α -helix. Even more significantly, the dispersions based on pseudocontact shifts offer better chances for structural characterization of the dynamic species. This method can be generalized for a large class of applications via the use of specially designed lanthanide-binding tags.

Keywords Cardiac troponin $C \cdot Spin$ -lock relaxation dispersion experiment $\cdot \mu s$ -ms protein dynamics \cdot Lanthanide \cdot Pseudocontact shift \cdot Calmodulin

Introduction

Protein folding, protein–protein and protein–ligand interactions are all essentially dynamic processes that are often predicated on complex internal motions which occur on a μ s–ms time scale. For studies of μ s–ms dynamics a number of NMR experiments have been developed over the course of years (Palmer 2004; McDermott 2004). Notably, these experiments are uniquely suited for characterization of dynamic equilibria involving 'excited states', i.e. transient low-population species that often play a crucial role in the biological function of proteins (Mulder et al. 2001).

The data from NMR experiments are usually consistent with a simple two-state exchange model, where each of the states is characterized by distinct chemical shifts. The chemical shift difference between the two states, $\Delta\omega$, is central to NMR analyses of exchanging systems. Under fast exchange conditions, $\Delta\omega$ manifests itself through line broadening and can be accessed by means of relaxation dispersion experiments (Bloom



et al. 1965; Deverell et al. 1970). In this work we study the dynamic system where the difference in *chemical* shifts is augmented by the difference in *pseudocontact* shifts, induced by Ln³⁺ ions.

The proposed method is demonstrated for the N-terminal domain of cardiac troponin C. Troponin C (TnC) is a dumbbell-shaped protein which, together with troponin I (TnI) and T (TnT), forms the troponin assembly. The C-terminal lobe of TnC possesses two high-affinity metal-binding sites that are permanently occupied under physiological conditions. This domain plays a structural role as it anchors the N-terminal helix of TnI. In contrast, the N-terminal domain (NcTnC) serves as a Ca²⁺-regulated molecular switch. Only one of the two EF-hand motifs in NcTnC is capable of binding calcium—the other site is defunct due to alteration of the two ligand residues. Upon loading a single Ca²⁺ ion, NcTnC switches to the dynamic state that is primed for binding (Pääkkönen et al. 1998: McKay et al. 2000; Eichmüller and Skrynnikov 2005) and, consequently, locks onto a short amphiphilic helix III from TnI (Li et al. 1999). As a result, a large portion of TnI is displaced, including the two so-called inhibitory regions located up- and down-stream from the helix III (Farah et al. 1994; Takeda et al. 1997; Luo et al. 2000). These inhibitory regions are lifted off of their binding sites on the surface of actin, thereby bringing about changes in the actin-tropomyosin assembly and clearing the way for actin-myosin binding. The result is a contraction of a striated muscle (Kobayashi and Solaro 2005).

Like other calcium-binding proteins (Campbell et al. 1975; Lee and Sykes 1983; Bentrop et al. 1997), NcTnC binds Ln³⁺ ions in its calcium-binding site (Wang et al. 1981; Gay et al. 2004). Lanthanide ions have a well-established record as useful structural probes in NMR (Lee and Sykes 1983; Dwek et al. 1971), X-ray crystallography (Weis et al. 1991; Burling et al. 1996), and luminescence spectroscopy of proteins (Rhee et al. 1981). Particularly, in NMR studies contact and pseudocontact shifts, various relaxation enhancement parameters, and residual dipolar couplings due to Ln³⁺-induced alignment were all used for protein structure determination (Bertini et al. 2001a). In addition, Ln³⁺-for-Ca²⁺ substitution helps to elucidate the structural mechanism of calcium regulation (Mustafi et al. 2004; Dudev et al. 2005). In this work we extend the scope of Ln3+ applications by demonstrating that lanthanides can be used to 'highlight' the effects of µs-ms time-scale motion in proteins.

We have prepared the samples NcTnC with La³⁺, Ce³⁺, and Pr³⁺ ions and used our recent proton off-resonance spin lock experiment (Eichmüller and

Skrynnikov 2005) to record relaxation dispersion profiles. In the presence of paramagnetic lanthanides the dispersion amplitudes were strongly increased. We attribute this effect primarily to μ s time-scale dynamics in the lanthanide binding site. This dynamics leads to modulation of paramagnetic susceptibility tensor which, in turn, causes modulation of long-range pseudocontact shifts and thus gives rise to line broadening and dispersion effects.

We have also analyzed other possible forms of dynamics contributing to Ln^{3+} -induced dispersions. The data suggest that *opening/closing* motion such as identified in Ca^{2+} -NcTnC may well contribute to the observed dispersions. On the other hand, it is unlikely that *transient dimerization* of Ln^{3+} -NcTnC is directly responsible for the dispersion effects, although it may influence other forms of μ s motion. *On-off exchange* of Ln^{3+} ion does not seem to be a significant factor under the conditions of our experiments, and there is no reason to suggest a presence of secondary Ln^{3+} -binding sites.

In conclusion, we discuss the prospects for using ${\rm Ln}^{3+}$ ions as probes of native protein dynamics in the $\mu s-ms$ range. Remarkably, long-range pseudocontact shifts can generate large dispersions where chemical shifts fail because of their local nature. Even more significantly, the values of $\Delta \omega$ associated with pseudocontact shifts can be readily converted into structural parameters. This paves the way for the structural characterization of the exchanging species, which is especially valuable in the case of the elusive 'excited states'. However, this approach is feasible only if the ${\rm Ln}^{3+}$ ion is tightly bound in a well-defined static environment and does not interfere with native protein dynamics. From this perspective, lanthanide-binding tags present an attractive and sufficiently general option.

Observation and analysis of Ln3+-induced dispersions

The affinity of Ca^{2+} to NcTnC is ca. 3 μ M at 30°C (Li et al. 1997; Hazard et al. 1998) and probably somewhat tighter at lower temperatures (Li et al. 2002). The titration of Ca^{2+} , therefore, proceeds under intermediate-to-fast exchange conditions with each resonance moving in a continuous fashion. In contrast, the La^{3+} , Ce^{3+} and Pr^{3+} titration is characterized by a slow exchange regime, with apo and holo forms giving rise to distinct sets of peaks. In this situation, one cannot rely on a lanthanide titration to obtain complete spectral assignment of Ln^{3+} -NcTnC. Instead, a set of triple-resonance experiments was executed for each of the three samples and the backbone assignment was thus obtained. The ^{1}H , ^{15}N -HSQC spectra of the



NcTnC samples loaded with different ions are presented in Fig. 1.

The analysis of the lineshapes in the course of the lanthanide titrations established that the Ce³⁺ off-rates at 30°C are in the range of 15–25 s⁻¹, while Pr³⁺ off-rates at 10°C are 5–10 s⁻¹. Assuming that lanthanides bind to NcTnC in diffusion-controlled manner, just like calcium (Hazard et al. 1998; Li et al. 2002), one obtains $K_D \approx 0.1 \mu \mathrm{M}$, in agreement with previous results (Wang et al. 1981). This binding is sufficiently tight to avoid unwanted complications from on-off exchange (discussed in more detail later).¹

The spectra shown in Fig. 1 were used to determine pseudocontact shifts according to $\delta_{pc}(Ln^{3+})=\delta(Ln^{3+})$ $-\delta(La^{3+}),$ where Ln^{3+} is a paramagnetic ion (Ce $^{3+}$ or $Pr^{3+})$ and La^{3+} is diamagnetic lanthanum ion used as a reference. 2 The resulting δ_{pc} data were fitted to the well-known expression:

$$\delta_{\rm pc}^i = \frac{\Delta \chi_{\rm a}}{12\pi r_i^3} \left\{ (3\cos^2\theta_i - 1) + \frac{3}{2}R\sin^2\theta_i\cos2\phi_i \right\} \qquad (1)$$

where $\Delta \chi_a$ is the axial component of the (traceless) magnetic susceptibility tensor of the ${\rm Ln}^{3+}$ ion in its binding site, R is the rhombicity of this tensor, θ_i and ϕ_i describe the orientation of the vector connecting the ${\rm Ln}^{3+}$ ion with the spin of interest relative to the principal axes of the susceptibility tensor, and r_i is the corresponding distance (Bertini et al. 2001c). The total of 117 and 98 experimentally measured pseudocontact shifts (${\rm Ce}^{3+}$ and ${\rm Pr}^{3+}$ data sets, respectively) have been fitted with Eq. 1 using the NMR structure of ${\rm Ca}^{2+}$ -NcTnC 1AP4 (Spyracopoulos et al. 1997). The results of the fitting, Fig. 2, can be taken as a confirmation that the ${\rm Ln}^{3+}$ -for- ${\rm Ca}^{2+}$ substitution does not cause any major structural perturbations in the protein.

The degree of correlation observed in Fig. 2, r = 0.89, is somewhat lower than what is typically obtained for Ln³⁺-loaded proteins (Banci et al. 1996; Baig et al. 2004). For instance, for Dy³⁺-doped α -parvalbumin the correlation coefficient r = 0.95 has been obtained based on a similar fitting procedure (Baig et al. 2004). One

should bear in mind, however, that (*i*) Dy³⁺ generates much larger shifts than Ce³⁺ or Pr³⁺, (*ii*) in contrast to NcTnC, parvalbumin has very little internal motion (Baldellon et al. 1998), and, on a related note, (*iii*) the structural model of parvalbumin has somewhat higher precision. Of note, Ca²⁺-NcTnC also produced a relatively poor fit in the study employing ¹⁵N-¹H residual dipolar couplings (*r* around 0.8 for a number of structural models, including 1AP4) (Pääkkönen et al. 2000).

The sizeable pseudocontact shifts, up to 0.6 ppm, observed in Ce³⁺- and Pr³⁺-NcTnC spectra (Fig. 2) offer new possibilities for studies of us-ms time scale dynamics in this protein. Toward this goal, we recorded a large set of relaxation dispersion data in the Ln³⁺-loaded samples of NcTnC. The details of the experiment, developed with Ln³⁺-loaded proteins in mind, were recently reported (Eichmüller and Skrynnikov 2005). In designing this experiment we recognized that pseudocontact shifts, as expressed in ppm, do not depend on the spin's Larmor frequency. Therefore, in the case of chemical exchange the frequency modulation associated with pseudocontact shifts, $\Delta \omega = \omega_0 \Delta \delta_{pc}$, is much larger for protons than for heteronuclei. Thus, proton is the nucleus of choice for detecting Ln³⁺-induced dispersions (conversely, if the goal is to observe Ln³⁺-induced dispersions on heteronuclei, then conventional HSQCbased schemes should be abandoned because of the severe broadening in the proton dimension and directdetection schemes (Machonkin et al. 2002) should be employed instead).

The main difficulty in the proton relaxation dispersion experiments stems from extensive cross-relaxation and scalar coupling between numerous proton spins. To minimize these effects we implemented a number of solutions: (i) used deuterated samples with protons exchanged into amide positions (Ishima et al. 1998), (ii) employed off-resonance spin lock with the constant tilt angle of 35° (Desvaux et al. 1995), and (iii) utilized the spin mode with anti-phase character with respect to nitrogen, $(H_x + 2H_z)N_z$. Large effective rf fields obtainable in this experiment allow one to target faster forms of motion, down to several microseconds. Figure 3 shows the representative relaxation dispersion curves obtained from our constant-angle off-resonance ¹H^N spin lock experiment conducted on four holo-NcTnC samples. The relaxation rates determined in this experiment are, in a very good approximation, given by (Eichmüller and Skrynnikov 2005):

$$\bar{R}_{1\rho}^{\text{mod}} = R_{\text{ex}} + (R_{\text{anti}} - R_{zz}) \tag{2}$$

where R_{anti} and R_{zz} are the relaxation rates of H_xN_z and H_zN_z , respectively. The exchange contribution R_{ex}



 $^{^1}$ La³+, Ce³+, and Pr³+ have similar effective ionic radii, 1.10–1.06 Å, and therefore are all expected to bind in a similar fashion. Smaller ions, on the other hand, can behave differently. Indeed, we found that Yb³+ (0.92 Å) binds to NcTnC in the intermediate-to-fast exchange regime.

 $^{^2}$ Note that paramagnetic lanthanide ions also produce weak alignment (Contreras et al. 1999; Biekofsky et al. 1999) so that the experimentally measured differences $\delta({\rm Ln}^{3+})-\delta({\rm La}^{3+})$ contain, generally speaking, a residual CSA component (John et al. 2005). It is easy to estimate, however, that with the present very small degree of alignment the residual CSA shifts do not exceed 0.01 ppm and therefore can be safely ignored.

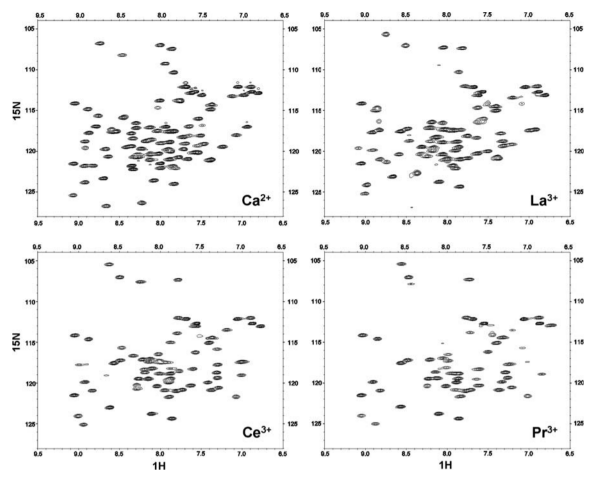


Fig. 1 ¹H, ¹⁵N HSQC spectra of Ca²⁺-, La³⁺-, Ce³⁺-, and Pr³⁺-NcTnC at 10°C, 600 MHz. The samples contained 1.2, 1.2, 2.0, and 2.0 mM of protein, respectively. The measurement times were adjusted to compensate for the difference in concentration; the spectra were plotted at the same contour level. Metal ion

content was 4.0, 1.25, 1.25, and 1.25 molar equivalents, respectively. In the case of Ca²⁺, the peaks from Gly 70 and Val 72 are outside the plotted spectral region. The tables of spectral frequencies for Ln³⁺-NcTnC (C35S, C84S) are given in the Supplementary Materials

for the system in question falls in the Redfield limit; in a two-state approximation, this term is expressed as (Deverell et al. 1970):

$$R_{\rm ex} = p_{\rm a} p_{\rm b} (\omega_0 \Delta \delta)^2 \frac{\tau_{\rm ex}}{1 + (\omega_1^{\rm eff} \tau_{\rm ex})^2}$$
 (3)

The dependence of the recorded dispersion profiles on the static magnetic field confirms that the observed exchange is fast (see Fig. 3) (Millet et al. 2000).

The curves presented in Fig. 3 illustrate two types of behavior. Residue 14 shows no dispersion in any of the samples. Apparently, there is no local motion that could modulate the ¹H^N chemical shift in this residue. This residue is also located too far from the metal center (Ce³⁺, Pr³⁺) to sense any significant pseudocontact shift. The situation is different in residue 42. This residue shows no discernible dispersion in the diamagnetic Ca²⁺-NcTnC and La³⁺-NcTnC samples.

The dispersion appears, however, in the sample containing a weak paramagnetic ion, Ce³⁺-NcTnC. It is amplified in the presence of the stronger paramagnetic ion, Pr³⁺-NcTnC. This kind of behavior unequivocally suggests that the observed dispersions are caused by modulation of the pseudocontact shifts.

A systematic comparison of the dispersion profiles, beyond the two examples shown in Fig. 3, reveals a more complex pattern. Nonetheless, the general trend is unmistakable: paramagnetic ${\rm Ln}^{3+}$ ions tend to magnify the dispersion effects in this protein. This is demonstrated in Fig. 4 where the fitted values of $p_a p_b (\Delta \delta)^2$ obtained from the analyses of the relaxation dispersion data are plotted as a function of residue number (the corresponding $\tau_{\rm ex}$ values are presented in Fig. 5).

First, compare the results from the two diamagnetic samples, Ca²⁺-NcTnC and La³⁺-NcTnC (two top panels in Fig. 4). It is immediately obvious that there are substantial differences in the dynamic behavior of



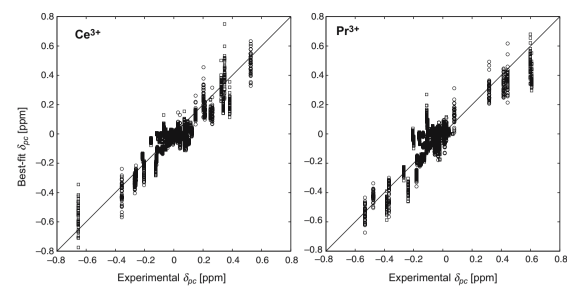


Fig. 2 The correlation between experimental and fitted pseudocontact shifts in Ce³⁺- and Pr³⁺-NcTnC. The experimental data are from ¹H, ¹⁵N-HSQC spectra recorded at 10°C at 600 MHz. Pseudocontact shifts of ¹H^N (circles) and ¹⁵N (squares) have been included in the fitting procedure with the same weight. Protein coordinates used in the fitting are from the PDB structure 1AP4 (Spyracopoulos et al. 1997), where Ca²⁺ has been replaced with Ln³⁺. A single susceptibility tensor has been determined in the course of the fitting procedure using all 40 conformations from 1AP4 as structural input (Banci et al. 1996; Eichmüller and Skrynnikov 2005). The best-fit parameters are $\Delta_{\chi_a} = 1.01 \cdot 10^{-26}$ ppm m³, R = 0.54, $\alpha = 33^{\circ}$, $\beta = 120^{\circ}$, $\gamma = 6^{\circ}$ for Ce³⁺-NcTnC and $\Delta_{\chi_a} = 1.65 \cdot 10^{-26}$ ppm m³, R = 0.57, $\alpha = 31^{\circ}$, $\beta = 126^{\circ}$, $\gamma = 2^{\circ}$ for Pr³⁺-NcTnC. The Euler angles α , β , γ specify

the orientation of the susceptibility tensor principal axes in the molecular frame 1AP4. The correlations presented in the plot are characterized by r=0.89, Q=0.47 and r=0.89, Q=0.46 for the left and right panels, respectively. In addition to 1AP4 we have also tested other sets of coordinates: NMR structure 1MXL (Li et al. 1999) (entire ensemble as well as individual conformations) and crystallographic structure 1J1D (Takeda et al. 2003). Furthermore, the fitting procedure was repeated using the reduced $\delta_{\rm pc}^i$ data set where we excluded the data from terminal helix D (likely to be affected by protein dimerization, see below). None of these modifications caused any significant change in the quality of data fitting or in the extracted susceptibility parameters

the two samples. The important feature of La³⁺-NcTnC is the mobility of the metal-binding EF-hand. Within the twelve-residue calcium-binding segment, the resonances from residues 70 and 72 are unobservable, the resonances from residues 66, 67, 69, and 73 are unobservable at 10°C but appear as weak and broadened peaks at 30°C, and residues 65, 68, 71, 75, and 76 show pronounced dispersion effects. The list of the affected residues includes all five of the canonical EF-hand ligands. This is in sharp contrast to Ca²⁺-NcTnC, where all twelve resonances are present and none of them shows statistically significant dispersion effects.

While at first this contrasting behavior seems surprising, it can be rationalized. NcTnC is a molecular switch with a finely tuned response to calcium binding (McKay et al. 2000). Although a good mimetic of calcium, La³⁺ carries an extra charge which is likely to perturb the delicate equilibrium existing in the Ca²⁺-loaded NcTnC. The result is 'dynamization' of the metal-binding site, residues 65–76. The changes are transmitted to the other parts of the protein via the network of hydrogen bonds, and the defunct calciumbinding site, residues 29–40, apparently responds to

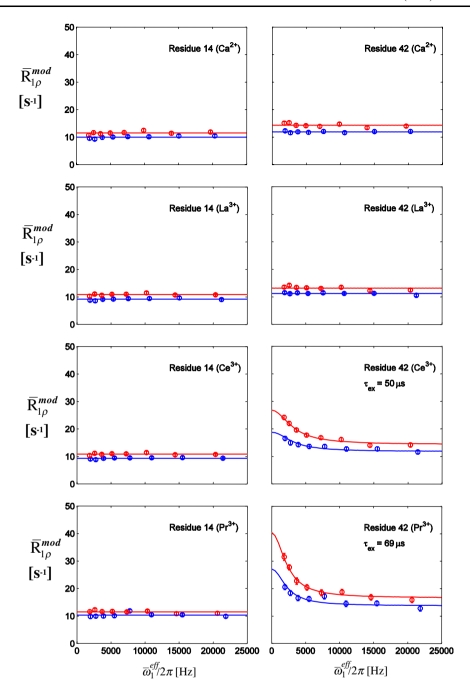
that by becoming more rigid. Of course, the disappearance of dispersions in this situation may also mean a shift in the time scale of motion which would render it unobservable for spin-lock measurements.

We shall turn now to the comparison of lanthanideloaded samples (three lower panels in Fig. 4). In the vicinity of the metal-binding site, residues 50-80, the comparison is hardly possible since most resonances in the Ce³⁺ and Pr³⁺ samples are broadened beyond detection. The major source of this broadening is a Curie spin relaxation (Guéron 1975; Vega and Fiat 1976) and hence no assumptions can be made about the effects of exchange. However, the remaining portion of the protein, residues 3-50, lends itself to comparison. The data in Fig. 4 clearly show that the amplitude of dispersion effects increases in going from diamagnetic La³⁺ to weakly paramagnetic Ce³⁺ and then to moderately paramagnetic Pr³⁺. This demonstrates, on a sufficiently large set of data, that modulation of pseudocontact shifts by exchange can be an effective mechanism of line broadening.

The important question that remains is: what type of molecular motion is responsible for modulation of



Fig. 3 Representative relaxation dispersion profiles obtained from Ca²⁺-, La³⁺-, Ce³⁺-, and Pr³⁺-NcTnC using constant-angle off-resonance proton spin-lock experiment. The data have been collected at 600 and 800 MHz (blue and red circles, respectively). The exchange correlation times indicated for Ce3+- and Pr³⁺-NcTnC have the error intervals 39–62 and 53–91 μ s, respectively. The details of the experiment, the fitting procedure, and the Ca²⁺-NcTnC results have been reported previously (Eichmüller and Skrynnikov 2005). The presented curves illustrate typical quality of the data. For example, if all data are ranked according to the fit residual, residue 42 consistently ranks in the 45-55 percentile



pseudocontact shifts? This question is addressed in the next section.

Possible dynamic scenarios for Ln³⁺-induced dispersions

Modulation of χ by internal motion in the Ln³⁺-binding site

We suggest that the primary cause of Ln³⁺-induced dispersions observed in NcTnC is the modulation of

the susceptibility tensor χ by μ s time-scale motion which occurs in the lanthanide-binding site. The modulation of χ automatically leads to modulation of pseudocontact shifts, cf. Eq. 1, which in turn leads to extensive dispersion effects. What is the evidence for this scenario? First of all, μ s time-scale dynamics in lanthanide-loaded EF hand is directly observed in the diamagnetic La³⁺-NcTnC sample (second panel in Fig. 4). As discussed above, the motion involves all lanthanide ligands. Note that even a slight rearrangement of the low-symmetry Ln³⁺ environment could be sufficient to alter the orientation of the χ principal axes



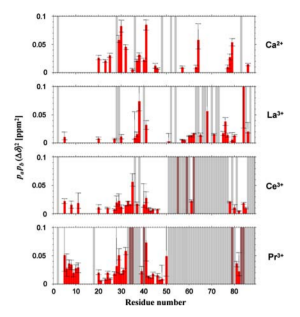


Fig. 4 Best-fit values of $p_ap_b(\Delta\delta)^2$ obtained from fitting of $^1H^N$ relaxation dispersion data in Ca^{2+} -, La^{3+} -, Ce^{3+} -, and Pr^{3+} -, NcTnC. Empty slots in the histogram indicate the absence of dispersion. Red bars represent $p_ap_b(\Delta\delta)^2$ values. Grey bars indicate missing data (this includes Pro 52 and Pro 54, a number of resonances that have been assigned at 30°C but disappear from the spectrum at 10°C, and several heavily overlapped peaks). Pink bars correspond to severely broadened peaks where dispersion is identifiable by eye, but is not picked up by the model selection criterion (Eichmüller and Skrynnikov 2005) because of the poor signal-to-noise ratio

and thereby cause effective modulation of the shifts. In this situation it is almost certain that χ undergoes efficient μ s time-scale modulation.

Let us assume for a moment that the μ s time-scale motion in the binding site can be viewed as a two-state exchange. The observed pseudocontact shifts in this case will be given by $p_a \delta_{pc}^a + p_b \delta_{pc}^b$, while the corresponding dispersion parameters will be $p_a p_b (\delta_{pc}^a - \delta_{pc}^b)^2$. Although there is no reason to expect a strong correlation between these two quantities, a certain pattern should exist—for instance, both quantities disappear for those $^1H^N$ spins that are far removed from the paramagnetic center. Indeed, out of 10 amide protons with the smallest pseudocontact shifts only 2 show dispersions, while among 10 protons with the largest shifts 9 show dispersions (Ce³⁺-NcTnC data; the results for Pr³⁺-NcTnC are similar). This is compatible with the proposed scenario whereby the dispersions are caused by modulation of χ .

In this context it is also worth noting that the values of $\Delta \chi_a$ determined in our analyses, Fig. 2, are two times lower than those found in another EF-hand protein, calbindin D_{9k} (Bertini et al. 2001b; Allegrozzi et al. 2002). In line with the evidence discussed above, this is suggestive of motional averaging of χ in Ln³⁺-NcTnC.

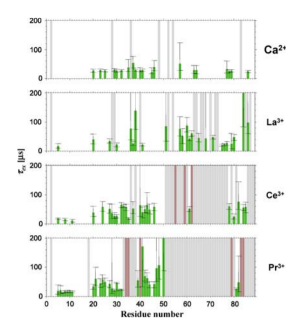


Fig. 5 Best-fit values of τ_{ex} obtained from fitting of ${}^{1}H^{N}$ relaxation dispersion data in Ca²⁺-, La³⁺-, Ce³⁺-, and Pr³⁺-Nc TnC. The conventions are the same as in Fig. 4

Within the framework of the present scenario it is also possible to rationalize the τ_{ex} data, Fig. 5. In the lanthanide-loaded samples, τ_{ex} values show some variability (in particular, shorter τ_{ex} are observed near the N-terminus). If one assumes that the motion in the Ln³⁺-binding site is actually more complex than two-state exchange, it is easy to see how different portions of the protein structure may sense somewhat different time scales.³ This kind of τ_{ex} variability is common for more complex forms of exchange in proteins (Mulder et al. 2001). Of note, there is much less variability in simple situations such as on-off exchange (Millet et al. 2002).

While no high-resolution coordinates are available for $\rm Ln^{3+}$ -NcTnC, inspection of calcium-loaded structures 1AP4 and 1MXL (Li et al. 1999) offers a hint as to the possible origins of μs time-scale dynamics in the binding site. The coordination of calcium by carboxylate oxygens of Asp 65 and Asp 67 shows substantial variability. In particular, the side chain of Asp 67

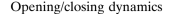
 $^{^3}$ For the sake of example, suppose that the binding site in Ln^{3+} -NcTnC adopts three different conformations, giving rise to pseudocontact shifts $\delta_{\text{pc}}^{\text{a}}$, $\delta_{\text{pc}}^{\text{b}}$, and $\delta_{\text{pc}}^{\text{c}}$. Further suppose that for a certain amide the orientation of the Ln^{3+} - $^1\text{H}^{\text{N}}$ vector is such that $\delta_{\text{pc}}^{\text{a}}$ and $\delta_{\text{pc}}^{\text{b}}$ are degenerate, whereas $\delta_{\text{pc}}^{\text{c}}$ takes a distinctive value. Under these circumstances, the dispersion profile will be sensitive to the exchange rates k_{ac} and k_{bc} , but essentially independent of k_{ab} . If the dispersion data are subsequently fitted with the two-state model (which usually produces a satisfactory fit) then the derived τ_{ex} will be representative of k_{ac} and k_{bc} , but not of k_{ab} . Reasoning along these lines, one can see how different $^1\text{H}^{\text{N}}$ sites may report somewhat different τ_{ex} values.



samples different conformations with regard to χ_1 , which directly affects the ligation of Ca^{2+} . Furthermore, in the structure 1MXL the side chain of Asp 65 turns out to be poorly localized with respect to Ca^{2+} . Occasionally, both aspartate side chains form hydrogen bonds with the backbone amide groups. In summary, the analysis of NMR coordinates suggests that there is room for dynamics in the metal-binding site of NcTnC.

In this context it is interesting to point out that μ s time-scale dynamics has been observed in the asparatic acid residue which ligates calcium in P43M mutant of calbindin D_{9k} (Bertini et al. 2002). Another wellcharacterized example of a dynamic binding site occurs in the Ca²⁺-saturated E140O mutant of the C-terminal domain of calmodulin (E140Q-CCaM). In this protein extensive motions have been observed not only in the binding site that is compromised by the mutation (129– 140), but also in the intact site (93–104) (Evenäs et al. 1999). A molecular dynamics study suggests that replacement of one Ca²⁺ ligand by a water molecule may be implicated in this process (Likic et al. 2003). The motion has been characterized as a two-state equilibrium involving open and closed conformations. The populations of the two species are approximately equal and the time scale of the exchange is ca. 20 µs (Evenäs et al. 1999; Lundström and Akke 2004). One cannot help but notice the similarity with Ca²⁺-NcTnC, where one of the calcium-binding sites is defunct, and the time scale of motion is 30 µs (Eichmüller and Skrynnikov 2005).

The possibility of conformational exchange has been also noted in the systems where Ln³⁺ is substituted for Ca²⁺. While the usual coordination number of Ln³⁺ in the EF-hand motif is seven, octa-coordinated Yb³⁺ has been found in the second site of carp parvalbumin (Kumar et al. 1991). The difference is due to Asp 92 which ligates Yb³⁺ via both carboxylate oxygens (this residue is equivalent of Asp 67 in NcTnC). The metalbinding site also proved to be flexible as indicated by the elevated temperature factors. In another study, Pidcock and Moore showed, by analyzing selected protein structures, that Ln³⁺-for-Ca²⁺ substitution tends to weaken the hydrogen bonding network around the metal-binding site (Pidcock and Moore 2001). While little information is available about proteins, in the case of small chelating complexes it is well known that bound ligands (including bidentate ligands) can rapidly interchange positions on the surface of the lanthanide ion and exchange with ligands that are not bound to the ion (Peters et al. 1996; Pidcock and Moore 2001; Parker et al. 2002). All this evidence supports the notion of dynamic lanthanide environment in Ln³⁺-NcTnC.



The most interesting possibility is that Ln³⁺-NcTnC retains much of the internal dynamics that occurs in Ca²⁺-NcTnC. If that is the case, then Ln³⁺-induced dispersion should reflect this 'native' dynamics—notably, the opening/closing motion proposed for Ca²⁺-NcTnC. This possibility is discussed below in detail.

Calmodulin-family proteins rely on a pair of sequential helix-loop-helix motifs to execute their function. In the apo protein, the two motifs are in UU-configuration, which corresponds to a closed form. In response to calcium binding, they switch to a more open L J conformation. The opening exposes a hydrophobic surface and thus allows proteins to bind their molecular targets.

NcTnC does not quite conform to the usual rules. Only one of the two EF-hands in NcTnC binds calcium, and the binding fails to achieve complete opening of the structure (Spyracopoulos et al. 1997). Instead, it is believed that Ca2+-NcTnC establishes dynamic equilibrium between the closed and open forms, with the closed form prevailing (Pääkkönen et al. 1998; McKay et al. 2000; Gaponenko et al. 1999). The protein can be locked in the open form when presented with a suitable molecular target, such as troponin I (Li et al. 1999) or bepridil (Li et al. 2000). The transition from the closed to open structure involves reorientation of helices B and C which move as a single unit in the direction away from the metalbinding site. Previously, this type of opening/closing dynamics has been identified in (Ca²⁺)₂-E140Q-CCaM (Evenäs et al. 1999) and proposed for apo-CCaM (Malmendal et al. 1999; Tjandra et al. 1995).

The concept of dynamic equilibrium involving closed and open forms of Ca^{2+} -NcTnC is consistent with our relaxation dispersion data (Fig. 4, upper panel). This is illustrated in Fig. 6. The left part of the figure shows dynamic residues, as identified in our relaxation dispersion measurements (Eichmüller and Skrynnikov, 2005), mapped onto the structure of the protein. The right part shows the residues that undergo a large change in dihedral angles ϕ , ψ upon opening of the protein structure (Spyracopoulos et al. 1997; Li et al. 1999). The correspondence between the two plots suggests that relaxation dispersions in Ca^{2+} -NcTnC arise, to a large extent, from the opening/closing of the molecular structure. Another source of dispersions is the inherent flexibility of the defunct Ca^{2+} -binding site.

Given the presence of opening/closing dynamics in Ca²⁺-NcTnC, it is possible that this type of motion is also responsible for Ln³⁺-induced dispersion effects. To test this hypothesis, we simulated the expected



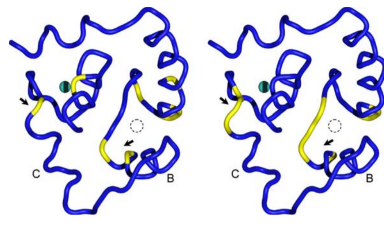


Fig. 6 Opening/closing in Ca²⁺-NcTnC. Left part: color-coded yellow are the residues with substantial dispersion amplitude, $p_{\rm a}p_{\rm b}(\Delta\delta)^2\!>\!0.03~{\rm ppm}^2$ (see upper panel in Fig. 4). Right part: color-coded yellow are the residues with $(\Delta\phi^2+\Delta\psi^2)^{1/2}>40^\circ,$ where $\Delta\phi$ and $\Delta\psi$ are the differences in the mean backbone dihedral angles between the closed structure 1AP4 and the open

structure 1MXL. The wireframe representation is that of the mean structure 1AP4 (minus flexible termini). Cyan sphere is the Ca²⁺ ion, dashed circle marks the defunct binding site, the arrows indicate the residues that play the role of hinges in the opening/closing of Ca²⁺-NcTnC, symbols B and C denote the corresponding helices

dispersion effects. We started by re-fitting the susceptibility tensor based on a reduced experimental data set. Excluded were all $\delta_{\rm pc}$ data from helices B and C (residues 38–64). The retained portion of the protein, which includes the metal-binding site, is deemed static and insensitive to the conformational exchange (in doing so we ignore the internal motions discussed in the previous section). The fitted susceptibility tensor was used to predict $\delta_{\rm pc}$ values in the closed structure 1AP4 and, separately, in the open structure 1MXL. The results were then subtracted to obtain $\Delta\delta_{\rm pc}$ which describes the presumed modulation of pseudocontact shifts by conformational exchange. Finally, the quantity $p_{\rm a}p_{\rm b}(\Delta\delta_{\rm pc})^2$ was evaluated, assuming that $p_{\rm a}$ is equal to 0.2 (McKay et al. 2000; Eichmüller and Skrynnikov 2005).

The simulated $p_a p_b (\Delta \delta_{pc})^2$ values plotted in Fig. 7 can be directly compared with the experimental results, two lower panels in Fig. 4. Such comparison shows that conformational exchange is likely to have a hand in Ln³⁺-induced dispersions for part of the defunct calcium-binding site and helix B (i.e. the region extending from residue 33 to residue 45). For instance, this model could explain the emergence of dispersion effect in residue 42, as illustrated in Fig. 3. This residue is a part of helix B which moves as a rigid unit when NcTnC undergoes transition from the closed to open form. Because the local environment of the ¹H^N from residue 42 does not change with motion, it shows no exchange broadening and no dispersion in the Ca²⁺and La³⁺-NcTnC samples. On the contrary, in the Ce³⁺- and Pr³⁺-NcTnC samples opening/closing leads to efficient modulation of δ_{pc} and hence gives rise to the dispersion effect.

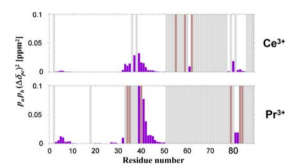


Fig. 7 Simulated $p_a p_b (\Delta \delta_{pc})^2$ values (purple bars) for the model of conformational exchange in Ce³⁺- and Pr³⁺-NcTnC. The format is the same as in Fig. 4. Open and closed forms of Ln³⁺-NcTnC were modeled by the NMR coordinates 1MXL and 1AP4, respectively. Prior to calculations, two structures were transformed into the same coordinate frame by superimposing their calcium-binding sites. The susceptibility tensors were determined using the invariant portion of the structure (δ_{pc} data from residues 38–64 were excluded, as described in the text). The best-fit parameters are $\Delta \chi_a = 1.14 \cdot 10^{-26}$ ppm m³, R = 0.64, α = 30°, β = 122°, γ = 1° for Ce³⁺-NcTnC and $\Delta \chi_a = 2.59 \cdot 10^{-26}$ ppm m³, R = 0.64, α = 35°, β = 123°, γ = 4° for Pr³⁺-NcTnC. In the latter case, $\Delta \chi_a$ is likely overestimated because the truncated data set contains only a few pseudocontact shifts that are not small. The above susceptibility parameters have been used to predict the values of δ_{pc} in 1AP4 and 1MXL and further calculate the difference, $\Delta \delta_{pc} = \delta_{pc}^a - \delta_{pc}^b$

Although this model can account for some of the experimental findings, it generally fails to predict strong dispersions observed in helices N and A (residue 3 to 28). In this situation it is difficult to make any claims with regard to observation of the exchange between open and closed conformations of NcTnC. It is almost certain that the competing mechanism—modulation of susceptibility tensor by local dynamics in the



binding site—is operational. It would be difficult to separate the two effects.

At the same time, one can take a view that the opening/closing in Ca²⁺-NcTnC and the fluxionality of the metal-binding site in Ln³⁺-NcTnC are nothing else than different manifestations of the vast dynamic network existing in this protein. This is discussed in more detail later.

Transient dimerization

It is known that weak protein self-association can cause exchange broadening (Pfuhl et al. 1999) and give rise to dispersion effects. Here we consider the feasibility of this scenario for Ln³⁺-NcTnC. In particular, we suggest that *intermolecular* pseudocontact shifts can come into play in this system, consistent with the increase in dispersion amplitudes along the series La³⁺-Ce³⁺-Pr³⁺.

EF-hand domains are known for their tendency to form dimers. The standard arrangement in the EFhand homodimers is such that the N-terminal helices from the two monomeric units pack against each other in antiparallel fashion, and likewise the C-terminal helices pack against each other in antiparallel fashion (Brodersen et al. 1998). The resulting pattern resembles a hash mark (#). A search of the Protein Data Bank (Berman et al. 2000) turned up 19 such structures (this number includes apo, Ca²⁺-loaded, and Ln³⁺loaded proteins, with structures solved both by X-ray (Mittl et al. 2002; Strynadka et al. 1997) and NMR spectroscopy (Potts et al. 1995; Drohat et al. 1999)). For Ca²⁺-NcTnC the evidence of weak self-association $(K_D = 7 \text{ mM})$ has been obtained from ¹⁵N linewidth measurements (Spyracopoulos et al. 2001).

To probe self-association effects in the lanthanide-loaded NcTnC we carried out the diffusion measurements using the ¹⁵N-filtered pulsed field gradient experiment (Choy et al. 2002). The data for a series of samples are illustrated in Fig. S1 and the results are summarized in Table 1. As can be seen from Table 1, the apo form of NcTnC shows substantial aggregation, which is somewhat alleviated when the sample is loaded with up to one molar equivalent of Ca²⁺ or Ln³⁺ ion. Subsequently, when the excess of an ion is introduced in the solution, the difference emerges between Ca²⁺ and Ln³⁺: the former shifts the equilibrium strongly toward the monomer, while the latter promotes aggregation.

The data listed in Table 1 have been used to estimate a fraction of dimers in solutions of Ln³⁺-NcTnC. As a first step, we employed HYDROPRO (de la Torre et al. 2000) to calculate the translational diffusion coefficients for the set of 19 EF-hand domain

Table 1 Translational diffusion coefficients in solutions of NcTnC

Loading ion content [molar equivalents NcTnC]	$D_t^{(a)}$ [10 ⁻⁷ cm ² /s]	Sample conditions ^(b)
0.0 1.0 (Ca ²⁺) 4.0 (Ca ²⁺) 4.0 (Ca ²⁺) 1.25 (La ³⁺) 1.25 (La ³⁺) 0.8 (Ce ³⁺) 1.25 (Ce ³⁺) 1.25 (Pr ³⁺) 1.25 (Pr ³⁺)	7.92 ± 0.03 8.15 ± 0.04 8.46 ± 0.10 (c) 8.61 ± 0.05 (d) 8.28 ± 0.03 7.82 ± 0.03 8.09 ± 0.05 7.84 ± 0.04 7.79 ± 0.07 8.37 ± 0.07	1.2 mM ² H, ¹⁵ N-NcTnC 1.3 mM ² H, ¹⁵ N-NcTnC 1.3 mM ² H, ¹⁵ N-NcTnC 1.3 mM ² H, ¹⁵ N-NcTnC; 100 mM NaCl ^(e) 0.15 mM ² H, ¹⁵ N-NcTnC 1.8 mM ² H, ¹⁵ N-NcTnC 0.9 mM ² H, ¹⁵ N-NcTnC 2.0 mM ¹³ C, ¹⁵ N-NcTnC 2.0 mM ¹³ C, ¹⁵ N-NcTnC 1.8 mM ² H, ¹⁵ N-NcTnC 1.8 mM ² H, ¹⁵ N-NcTnC; 100 mM NaCl

(a) The errors are determined by a jackknife method where two of the seven measured points (Fig. S1), are randomly discarded

(b) Other sample conditions are as described in the Materials and methods. Of note, addition of LnCl₃ aliquot produces turbid clouds in the solution of NcTnC. After approximately one minute of shaking, the sample returns to its original lucid state. This effect can be attributed to transient protein aggregation caused by a local excess of Ln³⁺ ions, consistent with the data presented in this table. Large excess of Ln³⁺ ions, above 2.5 molar equivalents, cause the protein to crash out of solution. None of this behavior has been observed in the Ca²⁺-loaded samples

(c) Assuming $D_t^{\text{monomer}} = 8.61 \cdot 10^{-7} \text{cm}^2/\text{s}$, $D_t^{\text{monomer}}/D_t^{\text{dimer}} = 1.20$ (see text) we obtain for this sample $K_D = 9$ mM. This is in good agreement with the previously reported result. $K_D = 7$ mM (Spyracopoulos et al. 2001)

(d) The calculation by HYDROPRO (de la Torre et al. 2000) using the structure of Ca²⁺-NcTnC (mean coordinates 1AP4) yields the value $D_t = 8.54 \cdot 10^{-7} \text{cm}^2/\text{s}$

(e) The effect of 100 mM NaCl on viscosity of the solution is negligible (Kestin et al. 1981)

homodimers found in the Protein Data Bank. This calculation led to the average ratio $D_t^{\rm monomer}/D_t^{\rm dimer}=1.20$. Using this ratio and assuming that $D_t^{\rm monomer}=8.61\cdot 10^{-7}{\rm cm}^2/{\rm s}$ (see Table 1) we estimated the fraction of dimers in each of our samples. For example, it has been found that the ${\rm Pr}^{3+}{\rm -NcTnC}$ sample in the absence of salt contains 55% of the dimeric form, whereas in the presence of salt the fraction of dimer drops to 15% (cf. two rows at the bottom of Table 1).

These estimates can be readily tested against the 15 N relaxation parameters. Assuming that $D_{\rm rot}^{\rm dimer}/D_{\rm rot}^{\rm monomer}=1.90$ (Daragan and Mayo 1997), we predicted that the addition of salt to the ${\rm Pr^{3+}}$ -NcTnC sample should sharpen the spectral lines by ca. 25%. Indeed, the addition of 100 mM NaCl caused uniform sharpening of the troponin resonances, with a 23% average reduction in the 15 N linewidth. Thus we are reasonably confident that our estimates correctly characterize the monomer–dimer equilibrium in lanthanide-loaded samples of NcTnC.



Certain tentative suggestions can be made about a structure of the Ln³⁺-NcTnC dimer. While NcTnC is distinct from most EF-hand domains in that it has an extra helix, its structure also permits a #-type arrangement found in the other EF-hand homodimers (illustrated in Fig. S2). Of note, this model predicts that the C-terminal residues from one monomeric unit are positioned near the metal-binding site of the other unit (and vice versa). This explains the disappearance of resonances 85-89 (86-89) in the samples loaded with Pr³⁺ (Ce³⁺), see Fig. 4. The peaks in question are likely obliterated by the intermolecular paramagnetic relaxation and by the Ln³⁺-induced exchange broadening. Apart from several C-terminal residues, the model shown in Fig. S2 does not predict any appreciable intermolecular contributions to pseudocontact shifts or to paramagnetic relaxation. This is also consistent with our observations and, in particular, with the fact that the experimental δ_{pc} data can be successfully fitted using the monomeric structure of NcTnC, Fig. 2.

The important question remains: is it possible that monomer–dimer exchange in the Ln³+-NcTnC samples is responsible for the observed relaxation dispersion behavior? To address this question let us assume that monomer and dimer are characterized by two distinct sets of shifts, δ^a and δ^b . In this situation the observed shifts, $p_a\delta^a+p_b\delta^b$, are a function of the monomer-dimer equilibrium. By utilizing the samples with different p_a/p_b , we can estimate $\Delta\delta=\delta^a-\delta^b$ and thus try to determine whether the monomer–dimer exchange can be a cause of the experimentally observed dispersions.

As it turns out, the change in p_a/p_b brings very little variation in the observed shifts. For instance, the addition of 100 mM NaCl to the Pr³⁺-NcTnC sample alters the monomer-dimer equilibrium from 45:55 to 85:15. At the same time, the ¹H^N spectral frequencies change only marginally (the largest shift 0.055 ppm, the second largest shift 0.035 ppm). Likewise, 10-fold dilution has little effect on the observed shifts: in dilute Pr³⁺ and La³⁺ samples the shifts change by less than 0.040 ppm compared to the concentrated samples. The pseudocontact shifts also remain essentially unchanged. These observations lead us to conclude that $\Delta \delta = \delta^a - \delta^b$ is small and, therefore, cannot be held responsible for the observed dispersion effects (in fact, our estimates show that $\Delta\delta$ falls an order of magnitude short).

This result is not so surprising from the structural standpoint. Indeed, it is often the case that formation of a dimer causes only limited changes in ¹H^N chemical shifts. We also do not expect to see much change in pseudocontact shifts, assuming that the structure of the dimer is as shown in Fig. S2.

The above analysis suggests that monomer-dimer exchange is unlikely to be the primary factor behind the observed dispersion effects. However, it is possible that dimerization affects the dispersion behavior indirectly. For instance, dimerization may alter the time scale of the internal motion in the Ln³⁺-binding site or the time scale of the opening/closing dynamics. To test this possibility, we recorded the relaxation dispersion data on the 100 mM NaCl Pr³⁺-NcTnC sample, where the dimerization is largely suppressed (600 MHz only, data not shown). Indeed, it turned out that the τ_{ex} in this sample are approximately 2-fold longer than in the sample that contains no salt. The details of weak self-association in Ca²⁺-NcTnC and the influence of this process on the exchange broadening are currently under investigation in Sykes' group (Julien and Sykes, personal communication).

Ln³⁺ on-off exchange

It has been demonstrated that the on-off exchange of the bound ion can give rise to relaxation dispersion effects in proteins (Millet et al. 2002). In the case of Ln³⁺-loaded samples, such exchange would automatically lead to modulation of pseudocontact shifts. In this section we argue that ion on-off exchange is unlikely to play any significant role with regard to the dispersion effects observed in the present study.

To determine the metal ion $k_{\rm off}$ rates, we analyzed the titration data from Ce³⁺ and Pr³⁺ samples (30 and 10°C, respectively). Specifically, in the series of ¹H,¹⁵N-HSQC spectra we selected several well resolved peak pairs from apo-NcTnC and Ln³⁺-NcTnC. For each of the apo peaks we monitored the broadening of the spectral line as a function of the lanthanide concentration. The linewidth data were fitted using the Bloch-McConnell equation (McConnell 1958), where the frequency separation between the apo- and holoresonances was taken directly from the spectra. The extracted off-rates were 5–10 and 15–25 s⁻¹at 10°C and 30°C, respectively.⁴

To minimize the effects of on-off exchange, the samples used for the dispersion measurements were

⁴ During the course of titration apo peaks behave exactly as expected, i.e. undergo gradual broadening to the point of complete disappearance. Holo peaks, on the other hand, do not show the expected sharpening trend. This unusual behavior can be explained if one recalls that the addition of Ln^{3+} promotes protein aggregation, which leads to longer τ_R and increased dipolar relaxation. Furthermore, the excess of Ln^{3+} brings about translational paramagnetic relaxation. The behavior of the holo peaks is therefore rather complex and does not lend itself to a quantitative interpretation.



loaded with 1.25 molar equivalents of La^{3+} , Ce^{3+} , and Pr^{3+} . The excess of lanthanide ensures that the fraction of apo-NcTnC is vanishingly small and, therefore, the exchange broadening is minimal. Using the determined value for $k_{\rm off}$, $10~{\rm s}^{-1}$ at $10^{\circ}{\rm C}$, and assuming that k_{on} is diffusion-controlled, as it is for calcium (Hazard et al. 1998; Li et al. 2002), we estimated that the $R_{\rm ex}$ contribution from the on-off exchange under these conditions does not exceed $0.2~{\rm s}^{-1}$. The small term of this magnitude can be safely ignored.

We can also approach this problem from another direction. Assume for a moment that the observed dispersion effects are due to the ${\rm Ln^{3+}}$ on-off exchange. Using the values of $\tau_{\rm ex}$ derived from the dispersion data (Fig. 5) and evaluating the relationships $\tau_{\rm ex}^{-1} = k_{\rm on}[{\rm Ln^{3+}}] + k_{\rm off}$ and $k_{\rm off}/k_{\rm on} = [{\rm NcTnC}][{\rm Ln^{3+}}]/[{\rm Ln^{3+}}] + {\rm NcTnC}]$ we can find $k_{\rm on}$ and the concentrations of apo and holo species in the sample. Based on these data, the $R_{\rm ex}$ contributions can be back-calculated. As it turns out, the $R_{\rm ex}$ values so obtained fall two orders of magnitude short of the experimentally measured values, which again demonstrates that the on-off exchange is not a principal cause of the dispersion effects.

As a final control, the dispersion measurements were repeated using an additional sample loaded with 0.8 molar equivalents of $\mathrm{Ce^{3+}}$ (600 MHz only, data not shown). In this sample, the contribution of on-off exchange to the transverse relaxation rate is expected to be approximately equal to k_{off} . The dispersion profiles recorded with this sample are in agreement, within error margin, with the main body of dispersion data.

On-off exchange involving secondary Ln³⁺ binding sites

In principle, it could be suggested that NcTnC has some additional low-affinity Ln3+ binding sites which could play a role in Ln³⁺-induced dispersions. There is, however, no factual basis for this hypothesis. In the case of Ca²⁺, no secondary binding has been observed in the thorough titration studies (Li et al. 1997). Likewise, Ce³⁺ and Pr³⁺ titration data contain no evidence of secondary binding. While some of the resonances drift slightly (up to 0.05 ppm in proton dimension) as the concentration of the lanthanide ion is raised from 1.0 to 1.5 molar equivalents, there are no dramatic shifts and all the peaks are accounted for throughout the titration. Finally, the control measurements conducted at 0.8 molar equivalents of Ce³⁺ confirmed that the presence of excess lanthanide ions has no impact on the observed dispersions.



The conformational state of the EF-hand pair is determined by three elements: two calcium-binding sites and the 2-helix element that straddles them. These elements are all structurally coupled. The sites are coupled with each other through a stretch of antiparallel β -sheet comprised of two residues (both residues are in position 8 of the canonical calcium-binding motif). They are also coupled to the 2-helix unit, with the C-terminal residue of site I (conserved Glu in position 12) and the N-terminal residue of site II (Asp or other residue in position 1) largely responsible for reorientation of this unit in response to ion binding (Grabarek 2006).

This assembly, termed ' $EF\beta$ -scaffold', displays substantial conformational diversity. For example, the orientation of the 2-helix element varies among different members of EF family, giving rise to the "conformational continuum" (Yap et al. 1999). Individual proteins also demonstrate conformational variability – for example, the N-terminal domain of calmodulin (NCaM) occurs not only in closed and open forms, but also in a semi-open form (Chou et al. 2001; Fallon and Quiocho 2003). A disulphide-linked variant of NCaM shows an unusual Ca^{2+} -binding mode where Glu in position 12 is not involved in coordinating the ion. It has been suggested that this conformation is also represented in the kinetic pathway of the native protein (Grabarek 2005).

With their predisposition to conformational heterogeneity, proteins such as CaM and TnC show the pattern of exchange broadening in the NMR spectra. The broadening generally occurs in and around the calcium-binding sites, but not in the 2-helix unit. In Ca²⁺-NcTnC, ¹⁵N line broadening has been previously noted for residues 28, 37, 39, 40, 61, and 64 (Pääkkönen et al. 1998), in agreement with our data. In apo-CCaM, ¹⁵N relaxation and relaxation dispersion experiments identified a number of residues affected by conformational exchange (Malmendal et al. 1999; Tjandra et al. 1995). If these data are mapped onto NcTnC (by aligning sites I of the two proteins), the residues 22, 23, 26, 28, 35, and 36 in and around the defunct binding site of NcTnC are predicted to be dynamic. In (Ca²⁺)₂-E140Q-CCaM, both binding sites show extensive broadening which has been linked to the opening/ closing motion (Evenäs et al. 2001; Lundström and Akke 2004). In particular, the largest exchange effects have been observed in the two β -sheet residues that form the bridge between the two calcium-binding sites.

Our data demonstrate an interesting complementarity effect in the μ s time-scale motion of the



calcium-binding loops. In the Ca²⁺-NcTnC sample, site I (defunct) is highly mobile, while site II (functional) is not, although the flanking residues immediately outside the calcium-binding loop show motion. In La³⁺-NcTnC the situation is reversed: the entire site II is mobile, but the motion in and around site I is considerably reduced. This is an interesting example of what can be called 'dynamic anti-cooperativity' in the pair of EF-hand motifs. From this perspective it is logical that the motions in Ca²⁺- and Ln³⁺-NcTnC occur on the same time scale (Fig. 5) as this time scale appears to be 'programmed' in the structure of the EF β -scaffold. Dynamic coupling between the two calcium-binding sites have been noted before in ¹⁵N relaxation studies of fast (ps-ns) motions (Akke et al. 1993; Spyracopoulos et al. 1998; Marchand and Roux 1998).

It is also noteworthy that the internal dynamics influences the affinity of the EF-hand site for the Ca²⁺ or Ln³⁺ ion. Based on the binding constants, the free energy difference between Ca²⁺- and La³⁺-NcTnC is estimated to be ca. 1–2 kcal/mol. The gain in enthalpy due to the electrostatic interactions is, however, much larger (Mustafi et al. 2004; Dudev et al. 2005). The balance is provided by entropy, which mainly stems from solvation/desolvation of the ions, but also reflects the change in internal dynamics of the protein.

Concluding remarks: Ln^{3+} ions as a probe of μ s-ms dynamics

Ideally, we would like to see Ln^{3+} as a non-intrusive probe of μ s-ms protein dynamics. In this desired scenario, Ln^{3+} binds tightly to a well-defined rigid binding site. In doing so, it provides a perfect substitute for calcium, preserving both the structure and dynamics of the physiologically-relevant Ca^{2+} -loaded form. When bound, Ln^{3+} induces pseudocontact shifts across a large swath of protein structure. If there is μ s-ms internal motion in the covered area, the involved spins experience modulation of pseudocontact shifts. As a result, the dispersion effects are amplified, or even created, for these spins.

In reality, however, we are faced with some contradictory choices. Among calcium-binding proteins we distinguish (i) the buffers and (ii) the sensor proteins. The buffers, such as parvalbumin, typically possess rigidly structured high-affinity binding sites (Corson et al. 1983), but feature little or none of μ s-ms dynamics (Baldellon et al. 1998). Conversely, sensor proteins, such as NcTnC, display rich and functionally important dynamic behavior. However, by their very nature they are finely tuned for regulation by calcium.

As a result, the Ln³⁺-for-Ca²⁺ substitution can be no longer considered perturbation-free. The assumption of a rigid binding site can also be called into question, as demonstrated in this study.

It is still feasible that this approach can work well for calcium-binding proteins. For instance, it can be envisaged that us-ms motions in multidomain proteins can be investigated in this manner by utilizing longrange pseudocontact shifts (Bertini et al. 2004). However, a more general approach is clearly desirable. We propose that the use of lanthanide-binding tags attached to the protein surface can provide a good solution. There are obvious advantages to this approach: (i) the Ln³⁺ probe can be introduced into proteins other than calcium-binding proteins, (ii) the tag can be positioned such that it provides good coverage of the dynamic site and at the same time does not interfere with the motion, (iii) the tags bind Ln³⁺ with high affinity and the coordination geometry is usually well characterized.

To explore this strategy we carried out some preliminary studies on NcTnC. A number of single-cysteine variants of NcTnC have been prepared, tagged with MTS-EDTA (Ikegami et al. 2004), and loaded with ions (with Ln³⁺ bound to a tag and Ca²⁺ bound to NcTnC). Of these mutants, S84C produced a poorly resolved spectrum suggestive of conformational heterogeneity. It is likely that in this case the tag interfered with protein opening/closing. On the contrary, T13C and D75C samples showed bona fide spectra. It turned out, however, that pseudocontact shifts in these spectra were small (less than 0.1 ppm for Yb3+-loaded tags). Furthermore, signal-to-noise ratio proved to be poor (presumably because of intermolecular Curie spin relaxation). As a result, these samples were unsuitable for observations of Ln³⁺-induced dispersions.

The main difficulty with the use of MTS-EDTA tags stems from the length and the flexibility of the linker. The 'swaying' motion of the tag is effective in averaging out pseudocontact shifts in the protein (Dvoretsky et al. 2002; Ikegami et al. 2004). Shorter and more constrained tags can produce larger pseudocontact shifts. Recently, a number of new tags have been developed that demonstrated good-quality spectra and pseudocontact shifts as large as 1 ppm (Ikegami et al. 2004; Prudêncio et al. 2004; Haberz et al. 2006). The use of these improved tags should facilitate the studies of Ln³+-induced dispersions.

Use of Ln³⁺ ions in relaxation dispersion studies offers some interesting possibilities. First, long-range pseudocontact shifts can generate large dispersions where chemical shifts fail. This happens, for example, when the motion involves a translocation of a 'rigid'



structural unit, such as α -helix. The local environment of the spins residing in such rigid fragment does not change and therefore there is no change in chemical shifts. Pseudocontact shifts, on the other hand, can be efficiently modulated when the fragment of interest moves relative to the Ln³⁺-binding site—thus generating the dispersion. The second and more important advantage stems from a simple relationship between pseudocontact shifts and structural variables. In the case of chemical shifts, it is generally difficult to translate $\Delta\omega$ values derived from the dispersions data into structural information (Hill et al. 2000; Skrynnikov et al. 2002; Korzhnev et al. 2004). In contrast, if $\Delta\omega$ originates from pseudocontact shifts then structural interpretation can be more straightforward. In conclusion, Ln³⁺-induced dispersions, demonstrated for the first time in this study, could become a useful addition to a growing collection of methods for studying µs-ms protein dynamics.

Materials and methods

The NcTnC (C35S/C84S variant) was expressed and purified as described previously (Li et al. 1995; Eichmüller and Skrynnikov 2005). The spectral assignment experiments were conducted on the samples of ¹³C, ¹⁵N-labeled NcTnC (1.2–2.0 mM protein concentration). The relaxation dispersion measurements were conducted on samples of ²H, ¹⁵N-labeled NcTnC (1.8 mM protein concentration). All samples were loaded with 1.25 molar equivalents of LnCl₃. The solvent composition was 90:10 H₂O:D₂O, 10 mM imidazole, 0.01% NaN₃, pH 7 (prior to Ln³⁺ titration). The S84C, T13C, and D75C mutants were constructed using the Stratagene QuikChange system. The tags were attached to cysteine thiol groups by incubating the samples overnight with MTS-EDTA (Toronto Research Chemicals). The presence of the Ln³⁺-loaded tag was confirmed by MALDI-TOF mass spectrometry. The sample conditions were 1.2-1.6 mM tagged protein, 1.0 molar equivalent Ln3+, 4.0 molar equivalents Ca²⁺, 150 mM NaCl, solvent as described above.

Backbone assignments of Ln³⁺-NcTnC were obtained from a standard suite of triple-resonance experiments including HNCACB, CBCA(CO)NH, HN(CA)CO, and HNCO (Kay 1995; Sattler et al. 1999) executed at 600 MHz Varian Unity Inova spectrometer at 30°C. The assignment was transferred to 10°C by means of the 'temperature titration' using ¹⁵N, ¹H-HSQC and HNCO experiments. The data were processed and analyzed using NMRPipe (Delaglio et al. 1995), NMRView (Johnson and Blevins 1994),

and Sparky (Goddard and Kneller 2002) software. The relaxation dispersion experiments and the data fitting procedure have been described previously (Eichmüller and Skrynnikov 2005). The dispersions data were collected at 12.4°C on Varian Unity Inova 600 and 800 MHz spectrometers equipped with room-temperature and cryogenic probes, respectively. The proton spin-lock field strengths varied from 1.13 (1.05) to 12.38 (11.75) kHz in the measurements at 600 (800) MHz. Each data set, collected in 27 h, was comprised of 6 reference spectra with $t_{rel} = 0$ and 16 spectra with $t_{\rm rel} = T = 50 \text{ ms}$ sampling 8 different spin lock strengths. The Ca²⁺-NcTnC data are as reported in our recent communication (Eichmüller and Skrynnikov 2005). Because of the data processing error, the 600 MHz dispersion profiles shown in Fig. 5 of that paper are shifted upward by a constant amount relative to their correct positions. This has strictly no impact on the analyses of slow motions as the shift is absorbed into $R_{\rm anti}^{600}-R_{zz}^{600}$. The curves shown in the top panels of Fig. 3 in this work are correct.

The translational diffusion data were obtained from a ¹⁵N-filtered pulsed field gradient experiment (Choy et al. 2002) executed in 1D mode. The measurement time was 3 h per sample. Gradient strengths were calibrated as described by Price (1998). The signals from the amide region were integrated using Varian VNMR software and the obtained integral intensities were analyzed as a function of gradient strengths as illustrated in Fig. S1. The HYDROPRO (de la Torre et al. 2000) calculations were performed assuming the temperature 10 °C and the corresponding solvent viscosity, 1.307 cP. In calculating the diffusion parameters we also used a randomly-selected control group consisting of 10 high-resolution structures of small dimeric proteins which produced the average value $D_t^{\text{monomer}}/D_t^{\text{dimer}} = 1.23.$

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