NMR Studies of Ca²⁺ Binding to the Regulatory Domains of Cardiac and E41A Skeletal Muscle Troponin C Reveal the Importance of Site I to Energetics of the Induced Structural Changes[†]

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ABSTRACT: Ca²⁺ binding to the N-domain of skeletal muscle troponin C (sNTnC) induces an "opening" of the structure [Gagné, S. M., et al. (1995) Nat. Struct. Biol. 2, 784-789], which is typical of Ca²⁺regulatory proteins. However, the recent structures of the E41A mutant of skeletal troponin C (E41A sNTnC) [Gagné, S. M., et al. (1997) Biochemistry 36, 4386-4392] and of cardiac muscle troponin C (cNTnC) [Sia, S. K., et al. (1997) J. Biol. Chem. 272, 18216-18221] reveal that both of these proteins remain essentially in the "closed" conformation in their Ca²⁺-saturated states. Both of these proteins are modified in Ca²⁺-binding site I, albeit differently, suggesting a critical role for this region in the coupling of Ca²⁺ binding to the induced structural change. To understand the mechanism and the energetics involved in the Ca²⁺-induced structural transition, Ca²⁺ binding to E41A sNTnC and to cNTnC have been investigated by using one-dimensional ¹H and two-dimensional ¹H, ¹⁵N}-HSQC NMR spectroscopy. Monitoring the chemical shift changes during Ca²⁺ titration of E41A sNTnC permits us to assign the order of stepwise binding as site II followed by site I and reveals that the mutation reduced the Ca²⁺ binding affinity of the site I by \sim 100-fold [from $K_{D2} = 16 \,\mu\text{M}$ [sNTnC; Li, M. X., et al. (1995) *Biochemistry* 34, 8330-8340] to 1.3 mM (E41A sNTnC)] and of the site II by \sim 10-fold [from $K_{\rm D1} = 1.7 \,\mu{\rm M}$ (sNTnC) to 15 μ M (E41A sNTnC)]. Ca²⁺ titration of cNTnC confirms that cNTnC binds only one Ca²⁺ with a determined dissociation constant K_D of 2.6 μ M. The Ca²⁺-induced chemical shift changes occur over the entire sequence in cNTnC, suggesting that the defunct site I is perturbed when site II binds Ca²⁺. These measurements allow us to dissect the mechanism and energetics of the Ca²⁺-induced structural changes.

Striated muscle contraction is initiated by Ca²⁺ binding to the thin filament protein TnC.¹ The resultant signal is transmitted to the other members of the thin filament (troponin I, troponin T, tropomyosin, and actin), which in turn modifies the interaction between the thick and thin filaments, leading to muscle contraction [for reviews, see Leavis and Gergely (1984), Zot and Potter (1987), Grabarek et al. (1992), Farah and Reinach (1995), and Tobacman (1996)]. Two isoforms of TnC exist in striated muscle, fast skeletal muscle TnC (sTnC) and slow skeletal/cardiac muscle TnC (cTnC). Both molecules comprise four putative EFhand helix—loop—helix motifs as potential Ca²⁺-binding sites (sites I—IV), except that site I in cTnC is inactive. Sites I and II are paired as a unit in the N-terminal half, and sites

III and IV form another pair in the C-terminal half of the molecule. It is generally believed that the role of paired sites III and IV is structural, whereas Ca²⁺ binding to sites I and II triggers muscle contraction [see reviews listed above; also see Potter and Gergely (1975) and Szczesna et al. (1996)]. While only site II in cTnC is essential for triggering cardiac muscle contraction, both sites I and II in sTnC are required for activating fast skeletal muscle (Putkey et al., 1989; Sheng et al., 1990).

The regulatory function of sTnC is associated with a structural change induced by Ca²⁺ binding to sites I and II in the N-domain. Although the crystal structure of avian sTnC was solved a decade ago (Herzberg & James, 1988; Satyshur et al., 1988), the Ca²⁺-induced structural transition of the N-domain remained undefined until the determination of the Ca²⁺-bound solution structures of the intact (Slupsky & Sykes, 1995) and N-domain of sTnC (Gagné et al., 1995) using NMR spectroscopy. The structures of the N-domain in both apo and Ca²⁺-saturated states (Gagné et al., 1995) define Ca²⁺-induced conformational transition from a "closed" state to an "open" state. This transition involves extensive exposure of a hydrophobic patch on the surface of the molecule long proposed as the interaction site for troponin I (Herzberg et al., 1986).

The regulatory function of cTnC is associated with structural change(s) induced by Ca²⁺ binding to site II in

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¹ Abbreviations: TnC, troponin C; sTnC, skeletal troponin C; cTnC, cardiac troponin C; cNTnC, N-domain (residues 1−89) of recombinant human cardiac troponin C; sNTnC, N-domain (residues 1−90) of recombinant chicken skeletal troponin C; NMR, nuclear magnetic resonance; HSQC, heteronuclear single-quantum coherence; NOE, nuclear Overhauser effect; $K_{\rm D1}$, dissociation constant of the first Ca²+ binding; $K_{\rm D2}$, dissociation constant of the second Ca²+ binding.

the N-domain. We have recently solved the solution structure of intact cTnC in the Ca²⁺-saturated state (Sia et al., 1997) and the structures of the regulatory N-domain in both apo and Ca²⁺-saturated states (Spyracopoulos et al., 1997). Surprisingly, it was found that unlike in sNTnC, cNTnC remains essentially closed in the Ca²⁺-saturated state, as a consequence of the defunct site I in cTnC. We have also recently solved the solution structure of a mutant of the N-domain of sTnC (E41A sNTnC) in which the bidentate Ca²⁺ ligand Glu 41 in site I is replaced by a nonliganding residue (Gagné et al., 1997). The structure of E41A sNTnC also remains closed upon Ca²⁺ binding. Both the E41A mutant and cTnC are modified in site I, albeit differently, suggesting a critical role for this region in the direct linkage between Ca²⁺ binding and the opening of the regulatory domain.

 Ca^{2+} ion chelation in a typical EF-hand involves seven liganding positions arranged as a pentagonal bipyramid. The six amino acid residues contributing to these liganding groups are designated in the sequences as positions X, Y, Z, -Y, -X, and -Z, respectively [see Strynadka and James (1989)]. In the site I of sNTnC, the Asp 30 in X, the Asp 32 in Y, and the bidentate Glu 41 in the -Z position are important ligands (Strynadka et al., 1997). In cNTnC, these two aspartic acid residues are replaced by Leu and Ala residues with no Ca^{2+} -coordinating side chains. These substitutions together with the insertion of a Val residue right before the loop I in cNTnC (Van Eerd & Takahashi, 1975) cause site I to lose its Ca^{2+} -binding ability.

Although these NMR solution structures provide vital information on the end states (apo and Ca²⁺-bound), it is important to understand the mechanism and energetics of the direct coupling between Ca2+ binding and induced structural changes in these regulatory domains. Previously, we have studied the Ca²⁺ binding mechanism of sNTnC by using two-dimensional (2D) {1H,15N}-HMQC NMR spectroscopy (Li et al., 1995). This approach was also applied by others to investigate Ca²⁺ binding to calbindin D_{9k} (Wimberly et al., 1995; Linse & Chanzin, 1995) and to calmodulin (Evenäs et al., 1997) and has been proven to be powerful because it can reveal information that pertains to individual atoms throughout the protein sequence. Our results demonstrated that Ca2+ binding to sNTnC occurs in a stepwise manner with the Ca2+ affinity of one site being approximately 10-fold stronger than that of the other. In this work, we used this approach to study Ca²⁺ binding to E41A sNTnC and cNTnC. During the processes of solving the NMR solution structures of E41A sNTnC and cNTnC, we have completely assigned the 2D {1H,15N}-HSOC spectra of E41A sNTnC and cNTnC in both the apo and Ca2+saturated states. These assignments are used in this paper to monitor the detailed Ca²⁺ titrations of E41A sNTnC and cNTnC. The results are used to evaluate the effect of the E41A mutation on the Ca²⁺ affinities of sites I and site II in sNTnC and on the Ca²⁺ binding energetics. Comparison of the Ca²⁺ binding properties of E41A sNTnC and cNTnC allows us to understand further the differences between slow/ cardiac and fast skeletal muscle contraction.

EXPERIMENTAL PROCEDURES

Nomenclature of Proteins and Mutants. The sequence numbering of chicken recombinant sTnC differs at two positions from that of human recombinant cTnC. The first residue in cTnC corresponds to residue 3 in sTnC, and there is a valine insertion at position 28 of the cTnC sequence. Following this numbering scheme, residue 90 in sTnC corresponds to residue 89 in cTnC. The designation sNTnC (1–90) is for the fragment encompassing residues 1–90 of chicken recombinant sTnC. E41A sNTnC (1–90) describes the same fragment in which Glu 41 is mutated to Ala. The designation cNTnC (1–89) is for the fragment encompassing residues 1–89 of human recombinant cTnC.

Construction of TnC Mutants. Protein Isolation, and NMR Sample Preparation. The engineering of the expression vector of E41A sNTnC (1-90) mutant was as described for sNTnC (Gagné et al., 1994), except the pTZ18·E41A sTnC plasmid was used as a template in the polymerase chain reaction (PCR) procedures. For the preparation of pTZ18. E41A sTnC, the 1.2 kb EcoRI DNA fragment carrying the sTnC gene was isolated from the pLcIIFx • sTnC (Reinach & Karlsson, 1988) and subcloned into the EcoRI site of pTZ18 plasmid DNA (Mead et al., 1986) to give pTZ18. sTnC. Clones carrying the insert in the right orientation were identified by single-stranded DNA sequencing. To isolate uracil-containing single-stranded DNA as the template for site-specific mutagenesis, pTZ18·sTnC plasmid DNA was transformed into Escherichia coli CJ 236 (ung⁻, dut⁻) cells (Kunkel et al., 1987). The nucleotide sequence of the mutagenesis primer was 5'-CACCAAGGCGTTGGGCA-3'. The underlined nucleotides were designed to substitute Ala for Glu at position 41 of sTnC. The mutagenesis mixture was transformed into E. coli JM109 (ung⁺, dut⁺) cells. Single-stranded DNA from several clones was isolated, and the E41A mutation was confirmed by DNA sequencing. The engineering of the expression vector of cNTnC (1-89) was as described by Chandra et al. (1997).

The expressions of E41A sNTnC, [15N]E41A sNTnC, cNTnC, and [15N]cNTnC in E. coli were as described previously for sNTnC and [15N]sNTnC (Gagné et al., 1994; Li et al., 1995). Purification of the proteins followed the previously published procedure for cleaved TnC (Golosinska et al., 1991). Decalcification of E41A sNTnC, [15N]E41A sNTnC, cNTnC, and [15N]cNTnC and their NMR titration sample preparations were as described for sNTnC (Li et al., 1995). All the NMR samples were in a 500 μ L volume. The buffer conditions were 100 mM KCl and 10 mM imidazole in 90% H₂O/10% D₂O for all the E41A sNTnC NMR samples and were the same for all the cNTnC NMR samples except there was 15 mM dithiothreitol (DTT) in the buffer; the pH was 6.70. The concentrations of [15N]E41A sNTnC, [15N]cNTnC, and cNTnC NMR titration samples were determined to be 2.08, 1.60, and 2.50 mM, respectively, by amino acid analyses. The concentration of the [15N]E41A sNTnC sample used for T2 measurements was determined to be 1.16 mM by amino acid analyses.

 Ca^{2+} Titrations of [15 N]E41A sNTnC, cNTnC, and [15 N]-cNTnC Monitored by 1D and 2D { 1 H, 15 N}-HSQC Spectra. A stock standardized 50 mM CaCl₂ solution and a stock 1 M CaCl₂ solution in water were used for the titrations. A 10 μ L Hamilton syringe was used for all the CaCl₂ solution additions. For the titration of [15 N]E41A sNTnC, aliquots of 2, 3, 3, 3, 4, 4, 4, 4, 5, 5, 5, 8, and 10 μ L of 50 mM CaCl₂ were added consecutively to the NMR tube for 14 individual titration points, and a 3 μ L sample of 1 M CaCl₂ was added for the last titration point. The total volume

changed from 500 μ L before titration to 566 μ L after titration. Both 1D and 2D { 1 H, 15 N}-HSQC spectra were acquired at every titration point. For the titration of [15 N]cNTnC, aliquots of 2, 3, 3, 3, 4, 4, 4, 4, 5, and 10 μ L of 50 mM CaCl₂ were added consecutively to the NMR tube for 10 individual titration points. The total volume increase was 42 μ L. Only 2D { 1 H, 15 N}-HSQC spectra were acquired during this titration. We performed 1D NMR titration of cNTnC separately by adding aliquots of 2, 2, 3, 3, 3, 3, 4, 5, 5, 5, 5, 10, 10, 10, and 10 μ L of a 50 mM CaCl₂ solution consecutively to the cNTnC NMR sample with a total volume change of 80 μ L. The changes in protein and Ca²⁺ concentrations due to dilutions were taken into account for data analyses. Changes in pH associated with CaCl₂ additions were negligible.

 T_2 Measurements As Monitored by 1D NMR Spectra. A jump and return spin echo pulse sequence (Anglister et al., 1993) was used to determine the effective rotational correlation time $\tau_{\rm ROT}$ for the protein E41A sNTnC. $\tau_{\rm ROT}$ is proportional to the molecular weight; $\tau_{\rm ROT} = (^4/_3\pi r^3)(\eta/kT)$. To a 500 μ L sample of E41A sNTnC were added consecutively aliquots of 1, 1, 1, 1, 1, 1, 1, 2, 2, 2, 2, 2, 2, 2, 2, 2, and 2 μ L of 50 mM CaCl₂. At every point, the T_2 was measured for the envelope of the NH resonances by using $1/T_2 = \pi \Delta \nu$ and $\tau_{\rm ROT}$ calculated from $\tau_{\rm ROT} = ^1/_5 T_2$ (Anglister et al., 1993). The temperature was 30 °C.

NMR Spectroscopy. All 1D and 2D {1H,15N}-HSQC spectra were recorded on a Varian VXR-500 NMR spectrometer. The 1D spectra were acquired using a spectral width of 7000 Hz, a pulse width of 9.8 μ s (90°), and an acquisition time of 2 s. The 2D {1H,15N}-HSQC spectra were acquired using the modified pulse scheme of Bodenhausen and Reuben (1980) with spin lock water suppression as described in Messerle et al. (1989). The temperature for all the experiments was 30 °C. Each 2D spectrum was collected with 512 complex data points in the t_2 dimension and 64 complex data points in t_1 ; eight transients per FID were recorded. The ¹H and ¹⁵N sweep widths were 7000 and 1500 Hz, respectively. The 2D {1H,15N}-HSQC spectra were processed using the software package NMRPipe (Delaglio et al., 1995) and peak-picked using the program PIPP (Garrett et al., 1991).

RESULTS

Ca²⁺ Titration of E41A sNTnC. A series of 1D ¹H NMR spectra, depicting the 8.8-11 ppm region of the spectrum, during the Ca²⁺ titration of [15N]E41A sNTnC are shown in Figure 1A. The two resonances near 10.3 ppm correspond to G35 and G71 (Gagné et al., 1997). The resonances for both amide NH protons appear as doublets due to coupling to the ¹⁵N. The chemical shift of these resonances changes as a function of the fraction of saturation of the protein with Ca²⁺ (Figure 2). In addition, these signals start as sharp resonances, undergo a slight line broadening, and then sharpen as the [Ca]_{total}/[E41A sNTnC]_{total} ratio is increased, indicating that the Ca²⁺ exchange kinetics fall between the intermediate and fast exchange limit on the NMR time scale (Li et al., 1995). The distinct shape of the Ca²⁺ binding plot (Figure 2A) for G35 clearly indicates the stepwise binding of Ca²⁺ to E41A sNTnC. The fact that the plot for G35 (Figure 2A) does not level off when the [Ca]total/[E41A sNTnC]_{total} ratio reaches 2 demonstrates that site I binds Ca²⁺

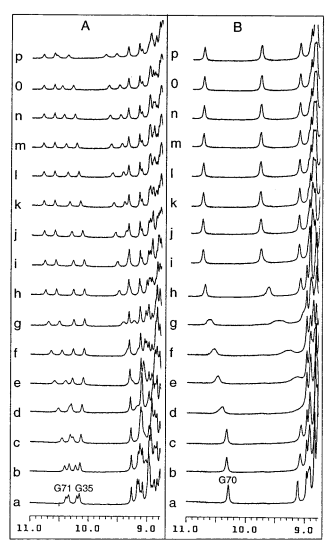


FIGURE 1: 500 MHz ¹H NMR spectra of 2.08 mM uniformly ¹⁵N-labeled E41A sNTnC (A) and 2.5 mM cNTnC (B), shown at [Ca]_{total}/[E41A sNTnC]_{total} ratios of (a) 0.000, (b) 0.051, (c) 0.208, (d) 0.376, (e) 0.526, (f) 0.677, (g) 0.885, (h) 1.094, (i) 1.302, (j) 1.511, (k) 1.771, (l) 2.037, (m) 2.298, (n) 2.715, (o) 3.235, and (p) 6.119 and [Ca]_{total}/[cNTnC]_{total} ratios of (a) 0.000, (b) 0.027, (c) 0.114, (d) 0.244, (e) 0.374, (f) 0.505, (g) 0.635, (h) 0.809, (i) 1.026, (j) 1.243, (k) 1.460, (l) 1.677, (m) 2.112, (n) 2.546, (o) 2.980, and (p) 3.415. Conditions are described in Experimental Procedures.

with a weaker affinity. On the other hand, G71 in site II is mainly affected by the binding of the first equivalent of Ca²⁺ (Figure 2B).

Virtually all residues of E41A sNTnC (except M1, D2, D3, T4, P53, and I73) can be observed in the 2D {\bar{1}H,\bar{1}SN}-HSQC spectra. The 2D {\bar{1}H,\bar{1}SN}-HSQC spectra acquired during Ca^{2+} titration were well-resolved at every titration point and have been completely assigned [see Gagné et al. (1997) and Figure S1 of the Supporting Information]. The cross-peaks shifted in the \bar{1}H and/or \bar{1}SN dimensions during Ca^{2+} titration. These chemical shift changes were plotted as a function of the [Ca]total/[E41A sNTnC]total ratio (see Figure S3 of the Supporting Information), as in the case of 1D titration. All of these Ca^{2+} binding curves were fitted individually to the equation

$$P + Ca \Leftrightarrow PCa + Ca \Leftrightarrow PCa_2$$

with K_{D1} and K_{D2} as the macroscopic dissociation constants for the binding of the first (site II) and second (site I) Ca^{2+}

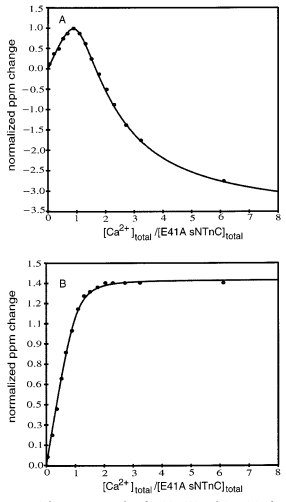
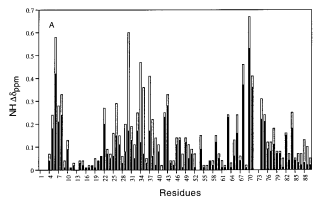


FIGURE 2: Calcium titration plots for G35 (A) and G71 (B) of E41A sNTnC derived from Figure 1A. The curve is normalized according to $(\delta_{\text{obs}} - \delta_{\text{initial}})/(\delta_{1\text{Ca}^{2+}} - \delta_{\text{initial}})$. The curves which best fit the data points are shown by the solid lines. The fitted results are shown in Results.

ions (Williams et al., 1985; Li et al., 1995). The fitting yielded a $K_{\rm D1}$ of 15 \pm 5 μ M and a $K_{\rm D2}$ of 1.3 \pm 0.5 mM. The resultant fitted curves for G35 and G71 are shown in Figure 2.

The plots of the absolute value of the difference of the amide ¹H and ¹⁵N chemical shifts for each residue between the apo and one-Ca²⁺, and one-Ca²⁺ and two-Ca²⁺ states, are shown in Figure 3. Chemical shift changes occur throughout the sequence when the [Ca]_{total}/[E41A sNTnC]_{total} ratio goes from 0 to 1, indicating that 1 equiv of Ca²⁺ is sufficient to induce global changes in this protein. Further chemical shift changes, also distributed throughout the sequence, occur when the second Ca2+ is added. This demonstrates that changes induced by Ca²⁺ binding to either site of E41A sNTnC are propagated globally. The most perturbed amides are those located in the two Ca²⁺ binding loops. These plots show clearly that loop II is more affected than loop I by binding the first Ca²⁺, whereas the second Ca²⁺ induces more changes in loop I than in loop II. Taken together, these results show that the E41A mutation reduced the Ca²⁺ binding affinity of site I in sNTnC by about 100fold ($K_{\rm D2} \approx 16~\mu{\rm M}$ to 1.3 \pm 0.5 mM, see Table 1) and of site II by about 10-fold (from $K_{\rm D1} \approx 1.7 \,\mu{\rm M}$ to $15 \pm 5 \,\mu{\rm M}$, see Table 1).



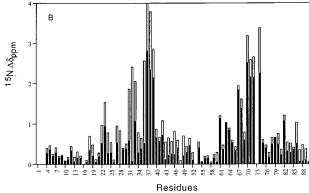


FIGURE 3: Plots of chemical shift changes (absolute values) for [¹⁵N]E41A sNTnC: (A) the apo to one-Ca²⁺ state (black bar) and additional changes after the one-Ca²⁺ state (open bar) for NH and (B) the apo to one-Ca²⁺ state (black bar) and additional changes after the one-Ca²⁺ state (open bar) for ¹⁵N.

Table 1: Ca^{2+} Binding Parameters for sNTnC, E41A sNTnC, and cNTnC^a

	$sNTnC^b$	E41A sNTnC	cNTnC
$K_{\rm D1}~(\mu{ m M})^c$	1.7 ± 1.0	15 ± 5	2.6 ± 0.1
$K_{\rm D2} (\mu {\rm M})^d$	16 ± 10	1300 ± 500	
ΔG°_{1} (kcal mol ⁻¹)	8.1	6.7	7.7
ΔG°_{2} (kcal mol ⁻¹)	6.9	4.1	

 a The binding constants were obtained as described in Results. The free energy was calculated from the standard relationships $\Delta G^{\circ}_{1} = RT$ ln $K_{\rm D1}$ and $\Delta G^{\circ}_{2} = RT$ ln $K_{\rm D2}$. b Data taken from Li et al. (1995). c Dissociation constants for the first Ca²⁺ bound (i.e., stronger site, which has been assigned to site II). d Dissociation constants for the second Ca²⁺ bound (i.e., weaker site, which has been assigned to site I).

Ca2+ Titration of cNTnC. Ca2+ titration of cNTnC was also monitored using both 1D ¹H and 2D { ¹H, ¹⁵N}-HSQC spectra. A series of 1D ¹H NMR spectra from the Ca²⁺ titration of cNTnC are shown in Figure 1B. The resonance at 10.3 ppm is G70 (Spyracopoulos et al., 1997), which is located in site II and corresponds to G71 in the sNTnC sequence. The chemical shift changes of this resonance are plotted as a function of the [Ca]total/[cNTnC]total ratio as shown in Figure 4. The plot saturates at a [Ca]total/ [cNTnC]_{total} ratio of 1, indicating that cNTnC binds only 1 equiv of Ca²⁺. The 2D {¹H, ¹⁵N}-HSQC spectra were very well-resolved, and the spectra in apo and Ca²⁺-saturated states are completely assigned [see Spyracopoulos et al. (1997) and Figure S2 of the Supporting Information]. All the residues along the cNTnC sequence except M1, D2, P52, and P54 were followed by ¹H and/or ¹⁵N chemical shift changes. All the resonances in the spectra were completely

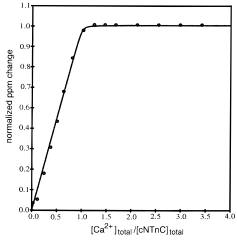
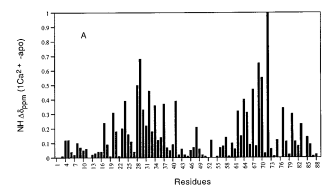


FIGURE 4: Calcium titration plot of G70 of cNTnC derived from Figure 1B. The curve is normalized according to $(\delta_{\text{obs}} - \delta_{\text{initial}})/(\delta_{1\text{Ca}^{2+}} - \delta_{\text{initial}})$. The curve which best fits the data points is shown by the solid line. The fitted results are shown in Results.



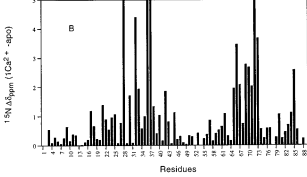


FIGURE 5: Plots of chemical shift changes (absolute values) for [15N]cNTnC: (A) the apo to one-Ca²⁺ state (black bar) for NH and (B) the apo to one-Ca²⁺ state (black bar) for ¹⁵N.

saturated when the [Ca]_{total}/[cNTnC]_{total} ratio reached 1. The data for all amides were fitted to the equation

$$P + Ca \Leftrightarrow PCa$$

(Williams et al., 1985) and yielded a macroscopic dissociation constant K_D of $2.6 \pm 0.1 \,\mu\text{M}$, which agrees with the Ca²⁺ affinity of site II in the native protein ($K_D = 1-5 \,\mu\text{M}$; Hannon et al., 1992; Johnson et al., 1980; Holroyde et al., 1980). Although only site II binds Ca²⁺, the chemical shift changes are distributed all along the sequence as shown in Figure 5, which plots the absolute chemical shift change differences for both amide ¹H and ¹⁵N for each residue. Figure 5 shows that most of the conformational perturbations by Ca²⁺ are located in the two Ca²⁺ binding loops and that, while site I is defunct, it responds to Ca²⁺ binding to site II.

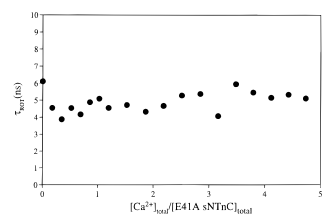


FIGURE 6: Plot of the effective rotation correlation time of E41A sNTnC determined from ¹H jump and return echo measurements of NH proton transverse relaxation at various [Ca]_{total}/[E41A sNTnC]_{total} ratios.

T₂ Measurements of E41A sNTnC. The rotational correlation time (τ_{ROT}) for the protein is proportional to its molecular weight, and also is influenced by the aggregation state of the protein. Figure 6 shows the τ_{ROT} values determined for E41A sNTnC during the Ca²⁺ titration, which vary in the range of 5.0 ± 1.1 ns corresponding to a molecular weight range of 7800-12200 with a median value of 10000, which is the molecular weight of E41A sNTnC (Sykes et al., 1996). This indicates that E41A sNTnC remains as a monomer upon Ca²⁺ binding in solution, unlike wild-type sNTnC (Gagné et al., 1995; Slupsky et al., 1995). The narrowness of the NMR line width and the resolution among the cross-peaks in the 2D {1H,15N}-HSQC NMR spectra during Ca²⁺ titration also indicate a lack of Ca²⁺induced aggregation/dimerization of E41A sNTnC in solution. The 2D {1H,15N}-HSQC spectra of [15N]cNTnC acquired during the Ca²⁺ titration are comparable to those of [15N]E41A sNTnC in terms of line width and resolution demonstrating no perceptible aggregation for cNTnC.

DISCUSSION

Activation of fast skeletal muscle involves binding of Ca²⁺ to two sites in the N-domain of sTnC, but only a single site in the N-domain of cTnC (Putkey et al., 1989; Sheng et al., 1990). To understand the differences in Ca²⁺ signaling between fast skeletal and cardiac muscle contraction, it is essential to have a thorough understanding of Ca²⁺ binding to the N-domains of the TnCs. We have previously studied in detail the mechanism of Ca²⁺ binding to the regulatory domain of wild-type skeletal TnC by 1D and 2D NMR spectroscopy (Li et al., 1995), and we have concluded that Ca²⁺ binding to the wild-type N-domain occurs in a stepwise manner with one site binding Ca²⁺ about 10 times stronger than the other; however, the data did not permit us to unambiguously assign the low- and high-affinity sites to structural sites I and/or II, respectively. On the basis of the amino acid residues occupying the liganding positions in site I and site II of sNTnC, we argued that site II is the site filled first. This conclusion has recently gained strong support from an analysis of the X-ray structure of sNTnC in the Ca²⁺bound state (Strynadka et al., 1997). Site I consists of residues **DADGGGDISTKE**, with six potential Ca²⁺coordinating liganding residues as bold letters. Positions 1 and 3 are two Asps coordinating directly to Ca²⁺, and position 12 consists of a Glu using both carboxylate oxygens

to coordinate to Ca²⁺; position 7 involves ligand formation with the main chain carbonyl group, and positions 5 and 9 involve water molecules coordinating to Ca²⁺. On the other hand, site II consists of residues DEDGSGTIDFEE. In comparison to site I, the Ser in position 5 interacts directly with Ca^{2+} through its O^{γ} atom. Thus, the main difference between the coordination of the Ca²⁺ ion is the fact that two water molecules are involved in site I but only one is involved in site II. In general, the more water ligands in the Ca²⁺ binding sites, the weaker the apparent Ca²⁺ affinity (Strynadka & James, 1989). In addition to the one more water molecule in site I, there is the presence of the conformationally flexible glycine triplet, the less overall negative charge for attracting Ca2+ ion, and the lower observed number of stabilizing hydrogen bonds (Strynadka et al., 1997). All these factors together would suggest that site I is energetically (both entropically and enthalpically) less favorable in binding Ca2+ than site II. This would explain that the greater Ca2+ binding free energy for site II $(\Delta G^{\circ} = -8.1 \text{ kcal mol}^{-1})$ with respect to site I $(\Delta G^{\circ} =$ $-6.9 \text{ kcal mol}^{-1}$) (see Table 1). By examining the Ca²⁺induced conformational changes in site II, Strynadka et al. (1997) have suggested that site II is more "pre-formed" for binding Ca²⁺ than site I in terms of ligand position and hydrogen bonding interactions, which would suggest that less conformational changes may be involved in Ca²⁺ binding to site II than to site I.

Gagné et al. (1997) have suggested that ligands in site I are essential in the coupling of Ca²⁺ binding to sites I and II in sNTnC to the induced structural change. In the E41A mutant, the bidentate ligand Glu 41 (position 12) in site I is replaced by an Ala. The structure of E41A sNTnC in the Ca²⁺-bound state remains closed, indicating that the direct linkage between calcium binding and the resultant opening of the structure is broken. On the basis of the new data reported here, we can now suggest that Ca2+ binding to sNTnC may involve the following steps. The first Ca²⁺ binds to site II, causing only minor structural changes and setting the stage for Ca²⁺ binding to site I, and then the second Ca²⁺ fills site I, inducing large conformational changes and the opening of the structure (Gagné et al., 1997). It should be noticed here that we have previously observed that Ca²⁺ binding to site II induced more significant far-UV CD ellipticity changes than binding to site I (Li et al., 1995), which, however, may not be directly related to the structural changes as discussed in Gagné et al. (1994). The opening of the structure and the exposure of the hydrophobic patch are energetically unfavorable in water. The energy barrier is estimated to be about 2.0 kcal mol⁻¹ by a recent lowtemperature, high-pressure study on sTnC (Foguel et al., 1996). In this paper, we found that site I is destabilized by $2.8 \text{ kcal mol}^{-1} \text{ (from } -6.9 \text{ to } -4.1 \text{ kcal mol}^{-1} \text{ , see Table}$ 1) with respect to Ca²⁺ binding due to the E41A mutation. This suggests that, when site I is altered by an E41A mutation, the reduction in Ca²⁺ binding free energy roughly compensates for the energetic barrier ($\Delta G^{\circ} = \sim 2.0$ kcal mol⁻¹) to open the structure and expose the hydrophobic surface. These measurements quantitatively support our conclusion (Gagné et al., 1997) that the opening of the sNTnC structure is dependent on one amino acid, Glu 41.

The present results also show that the E41A mutation destabilizes Ca^{2+} binding to site II by 1.4 kcal mol^{-1} (from -8.1 to -6.7 kcal mol^{-1} , see Table 1). Although

unexpected, this result is not surprising in light of the structural coupling between the two sites. The disposition of the two EF-hands and the central β -sheet connecting them in the N-domain of sTnC is characterized well in the crystal structures with the N-domain in apo states (Herzberg & James, 1988; Satyshur et al., 1988) and in the recent Ca²⁺bound structure of sNTnC (Strynadka et al., 1997). These studies suggest numerous hydrogen bonding and van der Waals interactions between the two sites, especially between the two β -strands connecting site I and site II. In the NMR structure studies of Ca²⁺-saturated sTnC (Slupsky & Sykes. 1995), sNTnC in apo and Ca²⁺-saturated state (Gagné et al., 1995), and E41A sNTnC in the Ca²⁺-bound state (Gagné et al., 1997), a large number of NOE contacts between the two sites were also observed. In the Ca²⁺ titration of sNTnC, we have observed that Ca²⁺ binding to either site II or site I causes chemical shift changes in both EF-hands (Li et al., 1995), suggesting widespread effects on conformational changes by each Ca²⁺ binding. In this work, we observed similar chemical shift changes for E41A sNTnC (Figure 3). In E41A sNTnC, the chemical shift changes are larger in site II for the binding of the first Ca²⁺ and larger in site I for the second Ca²⁺ clearly defining the order of the stepwise binding.

Mutating Glu to a nonbidentate ligand such as Gln or a nonligand such as Ala in the twelfth position of the consensus Ca²⁺ binding loop in other Ca²⁺ binding proteins (CaBP) has been carried out numerous times to investigate the critical role of this highly conserved residue for the Ca²⁺-induced conformational changes in the regulatory EF-hand proteins (Beckingham, 1991; Haiech et al., 1991; Maune et al., 1992; Carlström & Chazin, 1993; Evenäs et al., 1997). These mutations were observed to reduce drastically the affinity for Ca²⁺ in the mutated sites. Other site-specific mutagenesis studies have clearly defined these conserved Glu residues as an essential component for Ca²⁺ binding within the loops of TnC (Babu et al., 1992, 1993). The results of this study help to explain why this residue is highly conserved in the EF-hand Ca²⁺ binding proteins (Strynadka & James, 1989).

The E41A sNTnC structure stays closed in the Ca²⁺ state due to an imperfect Ca²⁺-binding site I in which the critical bidentate Glu 41 is replaced by an Ala. Interestingly, the cNTnC structure also remains essentially closed as a result of an inactive Ca²⁺-binding site I (Spyracopoulos et al., 1997). The 12-residue loop in the site I of cNTnC is LGAEDGCISTKE with liganding residues as bold letters. The important Glu 40 (position 12) is present, but the first two aspartic acid residues are replaced by residues (L and A) with no Ca²⁺-coordinating side chains, resulting in an inactive site I. In the case of E41A sNTnC, the substitution of Glu 41 by an Ala drastically reduced the Ca²⁺ binding affinity of this site but did not fully abolish the Ca²⁺ binding ability due to the effort of the remaining coordinating ligands (see above). In the case of cNTnC, the Glu 40 is present but the other ligands are not present so the Ca²⁺ cannot coordinate at all. There is therefore no Ca²⁺ for Glu 40 to interact with. The present results confirm the general belief [for a recent review, see Tobacman et al. (1996)] that there is no Ca²⁺ binding to site I. Our determined binding affinity for site II agrees well with literature values. The Ca²⁺ binding to site II of cNTnC yields ∼7.7 kcal mol⁻¹ (Table 1) of stabilizing free energy. The inactive site I in cNTnC also responds to Ca²⁺ binding to site II. This is due to the

structural coupling between the two sites. In the cTnC and cNTnC structures, we observed NOE contacts between the two loops, especially in the β -sheet. This result suggests that site I, though defunct, is not silent in the regulatory function of cTnC.

During the processes of solving the NMR structures of sTnC and sNTnC, we have noticed that these proteins dimerize and/or aggregate (Gagné et al., 1995; Slupsky et al., 1995). To assess the molecular state of E41A sNTnC in solution, we have determined the rotational correlation time (τ_{ROT}) for the protein at different [Ca]_{total}/[E41A sNTnC]_{total} ratios as shown in Figure 6. The results clearly indicate that E41A sNTnC does not undergo a Ca²⁺-induced dimerization/aggregation. This is consistent with the sharp NMR signals and the closed structure (Gagné et al., 1997). The sharp NMR signals and the observed closed structure for cNTnC suggest cNTnC also stays as a monomer in solution. Therefore, we conclude that our results presented in this paper are not influenced by protein dimerization/aggregation.

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SUPPORTING INFORMATION AVAILABLE

2D {¹H,¹5N}-HSQC spectra acquired during Ca²+ titration in Figure S1 for [¹5N]E41A sNTnC and in Figure S2 for [¹5N]cNTnC and Figure S3 showing additional examples of the Ca²+ titration curves as shown in Figure 2 (8 pages). Ordering information is given on any current masthead page.

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