Calcium-Dependent Stabilization of the Central Sequence Between Met<sup>76</sup> and Ser<sup>81</sup> in Vertebrate Calmodulin

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ABSTRACT Spin-label electron paramagnetic resonance (EPR) provides optimal resolution of dynamic and conformational heterogeneity on the nanosecond time-scale and was used to assess the structure of the sequence between Met<sup>76</sup> and Ser<sup>81</sup> in vertebrate calmodulin (CaM). Previous fluorescence resonance energy transfer and anisotropy measurements indicate that the opposing domains of CaM are structurally coupled and the interconnecting central sequence adopts conformationally distinct structures in the apo-form and following calcium activation. In contrast, NMR data suggest that the opposing domains of CaM undergo independent rotational dynamics and that the sequence between Met<sup>76</sup> and Ser<sup>81</sup> in the central sequence functions as a flexible linker that connects two structurally independent domains. However, these latter measurements also resolve weak internuclear interactions that suggest the formation of transient helical structures that are stable on the nanosecond time-scale within the sequence between Met<sup>76</sup> and Asp<sup>80</sup> in apo-CaM (H. Kuboniwa, N. Tjandra, S. Grzekiek, H. Ren, C. B. Klee, and A. Bax, 1995, Nat. Struct. Biol. 2:768–776). This reported conformational heterogeneity was resolved using site-directed mutagenesis and spin-label EPR, which detects two component spectra for 1-oxyl-2,2,5,5-tetramethylpyrroline-3-methyl)-methanethiosulfonate spin labels (MTSSL) bound to CaM mutants T79C and S81C that include a motionally restricted component. In comparison to MTSSL bound within stable helical regions, the fractional contribution of the immobilized component at these positions is enhanced upon the addition of small amounts of the helicogenic solvent trifluoroethanol (TFE). These results suggest that the immobilized component reflects the formation of stable secondary structures. Similar spectral changes are observed upon calcium activation, suggesting a calcium-dependent stabilization of the secondary structure. No corresponding changes are observed in either the solvent accessibility to molecular oxygen or the maximal hyperfine splitting. In contrast, more complex spectral changes in the line-shape and maximal hyperfine splitting are observed for spin labels bound to sites that undergo tertiary contact interactions. These results suggest that spin labels at solvent-exposed positions within the central sequence are primarily sensitive to backbone fluctuations and that either TFE or calcium binding stabilizes the secondary structure of the sequence between Met<sup>76</sup> and Ser<sup>81</sup> and modulates the structural coupling between the opposing domains of CaM.

INTRODUCTION
Calmodulin (CaM) is a ubiquitous eukaryotic calcium-binding protein that plays a critical role in intracellular signal transduction pathways (Crivici and Ikura, 1995; Tjandra et al., 1999). The crystal structure of calcium-saturated CaM indicates the presence of two globular domains joined by a solvent-exposed central helix located between Phe<sup>65</sup> and Phe<sup>92</sup> (Babu et al., 1985, 1988; Chattopdhyaya et al., 1992) (Fig. 1). Calcium activation of CaM, a prerequisite step for regulation of target proteins, involves a reorientation of the four α-helical bundles within each of the two globular domains with the concomitant exposure of hydrophobic sites that bind target proteins (La Porte et al., 1980; Kuboniwa et al., 1995; Urbauer et al., 1995; Zhang et al., 1995). Many of the structural details of this conformational transition have been documented. However, the presence of structural coupling between the opposing domains of CaM, in which one domain modifies the conformation of the other, and its functional role remains controversial.

Numerous biochemical studies indicate the presence of structural coupling between the opposing globular domains in CaM that may play an important role in modulating the kinetics of target protein activation (Yao et al., 1994; Pedigo and Shea, 1995a,b; Mukherjea et al., 1996; Shea et al., 1996; Sorenson and Shea, 1996; Sun et al., 1999; Jaren et al., 2000). For example, the proteolytic susceptibility or fluorescence intensity of sites in the amino-terminal domain responds to calcium occupancy to sites in the carboxy-terminal domain (Yao et al., 1994; Pedigo and Shea, 1995a; Shea et al., 1996, 2000; Sorenson and Shea, 1998). Similarly, the modification of sites within the carboxy-terminal domain (oxidation of either Met<sup>144</sup> or Met<sup>145</sup> or the site-directed substitution of Tyr<sup>138</sup> with Phe<sup>138</sup>) results in an enhanced rotational mobility of the amino-terminal domain (Gao et al., 1998; Yin et al., 2000a; Sun et al., 2001). There is, however, no consensus on the mechanisms underlying the observed structural linkages between the opposing domains, largely because multidimensional NMR measurements suggest that there are neither direct contact interactions between the opposing domains nor conformational coupling through the interdomain central sequence (Ikura et al., 1991; Barbato et al., 1992; Kuboniwa et al., 1995; Zhang et al., 1995). However, these latter NMR measurements selectively detect ordered structures that exhibit internuclear interactions and cannot resolve conformational...
heterogeneity on the nanosecond time-scale associated with the intramolecular reorientation of the opposing domains of CaM that is relevant to target protein binding.

Therefore, we have used site-directed spin-label electron paramagnetic resonance (EPR), which is sensitive to the nanosecond time-scale, to assess the dynamic structure of the sequence between Met\textsuperscript{76} and Ser\textsuperscript{81}. Five separate CaM mutants were created, permitting the incorporation of methanethiosulfonate spin-labels (MTSSL) covalently bound to individual cysteines engineered at defined positions within the primary sequence of CaM (Fig. 1). After spin-labeling, these sites permit a comparison of the dynamic structures of the sequence between Met\textsuperscript{76} and Ser\textsuperscript{81}. Five separate CaM mutants were created, permitting the incorporation of methanethiosulfonate spin-labels (MTSSL) covalently bound to individual cysteines engineered at defined positions within the primary sequence of CaM (Fig. 1). After spin-labeling, these sites permit a comparison of the dynamic structures of the sequence between Met\textsuperscript{76} and Ser\textsuperscript{81}, which have been suggested to be conformationally disordered compared with sites either in more stable regions of the central sequence or in globular domains expected to be sensitive to tertiary contact interactions (Finn et al., 1995; Mchaourab et al., 1996). Thus, individual cysteines were inserted at 1) T79C and S81C within the sequence between Met\textsuperscript{76} and Ser\textsuperscript{81}, 2) at F68C and L69C near the amino terminus of the central helix, and 3) at L105C in a helical site within the carboxyl-terminal globular domain.

We report that spin labels bound to either T79C or S81C, within the conformationally disordered sequence between Met\textsuperscript{76} and Ser\textsuperscript{81}, exhibit a spectral component with motional properties analogous to positions within stable helical regions within the central sequence (i.e., L69C) and that calcium activation stabilizes the motionally restricted conformation resulting in a more conformationally restricted and rigid interdomain sequence. This result is consistent with previous fluorescence measurements that demonstrated conformational changes involving the central sequence between Phe\textsuperscript{65} and Phe\textsuperscript{92}, which are important in the mechanism of calcium activation and target protein binding (Sun et al., 1999), and provides strong evidence that the opposing domains of CaM are structurally coupled through the interdomain sequence that includes the region between Met\textsuperscript{76} and Ser\textsuperscript{81}.

**MATERIALS AND METHODS**

**Materials**

1-Oxyl-2,2,5,5-tetramethylpyrroline-3-methyl-methanethiosulfonate spin label (MTSSL) was purchased from Reanal Fine Chemicals (Budapest, Hungary). WP PEI (weak anion exchanger) packing for the HPLC column was from J. T. Baker (Philipsburg, NJ), and the weak-anion exchange column used to purify CaM was packed in-house. A Micro BCA protein reagent kit was obtained from Pierce (Rockford, IL). HEPES was obtained from Research Organics, Inc (Cleveland, Ohio). All other reagent chemicals were of highest purity available commercially. Erythrocyte ghost plasma membranes were purified from porcine blood, essentially as previously described (Niggli et al., 1979).

**Expression and purification of CaM mutants**

The coding region for chicken CaM (accession number MCCH (PIR database) or P02593 (SWISS-PROT data base)) was excised from the plasmid pCaMPL provided by Professor Samuel George (Duke University) and subcloned into the mutagenesis and expression vector pALTER-Ex1 (Promega, Madison, WI) using restriction enzymes Ncol and Xbal. The recombinant plasmid pEx1-CaM was transformed into Escherichia coli strain JM109(DE3) for overexpression, as previously described (Studier and Moffatt, 1986; Sun et al., 1999). The sequence of chicken CaM is identical to all other expressed vertebrate CaM, including human, mouse, rat, rabbit, bovine, duck, frog, and salmon (Putkey et al., 1983; Wylie and Vanaman, 1988; Fischer et al., 1988; Kawasaki and Kretsinger, 1995; Friedberg, 1996). Furthermore, the measured mass of the expressed chicken CaM obtained using electrospray ionization mass spectrometry (ESI-MS) is 16,707 ± 3 Da, which is in good agreement with the theoretical mass of 16,706.4 for the following sequence: ADQLTIEQIA\textsuperscript{10} EFKEAFSLFD\textsuperscript{20} KDGDTITTK\textsuperscript{30} ELGTVMRSLG\textsuperscript{40} QNPTEALQD\textsuperscript{50} MINEVADAGN\textsuperscript{60} GITDPFELT\textsuperscript{70} MMARKKDTD\textsuperscript{80} SEEEIDREFR\textsuperscript{90} VFDKDGNYG\textsuperscript{100} SAAELRHVM\textsuperscript{110} LGELTDEE\textsuperscript{120} VDEMI- READ\textsuperscript{130} DGDQGVVNYEE\textsuperscript{140} FVQMIMTAK\textsuperscript{148}. This sequence contains no endogenous cysteines, permitting us to specifically introduce single cysteines for site-directed spin labeling. In all cases site-directed mutagenesis was carried out as described in the technical manual for the Altered Sites II in vitro mutagenesis system (Promega). Oligonucleotide primers containing the desired mutation were synthesized by Macromolecular Resources (Colorado State University, Ft. Collins, Co.). Correct mutations were ensured by automated DNA sequencing performed in the Biochemical Research Service Laboratory (University of Kansas, Lawrence, KS). Following overexpression, CaM was purified as previously described using phenyl Sepharose-CL-4B (Pharmacia, Piscataway, NJ) and Bakerbond WAX-HPLC (J. T. Baker, Phillipsburg, NJ) (Strasburg et al., 1988; Huhmer et al., 1996). The CaM concentration was determined using the Micro BCA assay (Pierce, Rockford, IL), using salt-free bovine brain CaM as a
standard. The concentration of the CaM standard was determined using the published molar extinction coefficient ($\epsilon_{280} = 3029 \text{ M}^{-1} \text{ cm}^{-1}$) for the calcium-saturated enzyme (Strasburg et al., 1988). The purity of the expressed proteins was greater than 99% as indicated by SDS-polyacrylamide gel electrophoresis and ESI-MS. Purified CaM was dialyzed against distilled water and following lyophilization was stored at $-70^\circ\text{C}$.

Covalent modification of CaM with MTSSL spin label

CaM (0.1 mM) was derivatized with a 10-fold molar excess of MTSSL in 25 mM HEPES (pH 7.5) and 1 mM EDTA at 25°C for $\sim$3 h, essentially as previously described (Berliner et al., 1982). The spin-labeled CaM was separated from unbound spin label using weak anion-exchange high-performance liquid chromatography (HPLC), essentially as previously described (Hühmer et al., 1996). Spin-labeled CaM was lyophilized to the desired volume and then reconstituted to activate the plasma membrane Ca-ATPase was assayed as previously described (Lanzetta et al., 1979; Yao et al., 1996).

Spectroscopic measurements

Circular dichroism (CD) spectra of CaM were measured at 25°C with a Jasco J-500CD spectrometer using a thermostatted CD spectral cell with a path-length of 1.0 cm. EPR spectra of 0.1 mM CaM were recorded in 50-μl disposable micropropettes (Rochester Scientific Co., Rochester, NY) in a Bruker ESP 300E X-band spectrometer (Billerica, MA) using a TM102 cavity fitted with a quartz dewar. Unless otherwise specified, spectra were acquired using an incident microwave field ($H_{\text{rf}}$) of 0.1 Gauss, a modulation frequency ($H_{\text{mod}}$) of 100 KHz, and a modulation amplitude ($H_{\text{amp}}$) of 2.0 Gauss. In all cases temperature was controlled to within 0.5°C using a Eurotherm B-VT 2000. Continuous-wave power saturation measurements of spin-labeled CaM were carried out essentially as previously described (Altenbach et al., 1989a; Mchaourab et al., 1996), where the power at which the signal amplitude is half that if no saturation occurred ($P_{1/2}$) was measured in the presence of oxygen saturated air ($P_{1/2}$) and nitrogen ($P_{1/2}$). The change in $P_{1/2}$ due to the presence of oxygen ($\Delta P_{1/2}$) is proportional to the collision frequency, where $\Delta P_{1/2} = P_{1/2}\text{(air)} - P_{1/2}\text{(N}_2)$. Additional terms that correct for spectrometer performance and differences in spin-lattice relaxation times permit a comparison between the solvent accessibilities of different spin-labeled CaM mutants, where the accessibility parameter ($\Pi$) is defined as:

$$
\Pi = \frac{\Delta P_{1/2}}{\Gamma(DPPH)} \times \frac{\Gamma(DPPH)}{\Gamma(DPPH)}
$$

where $P_{1/2}$ is the $P_{1/2}$ for a DPPH crystal, $\Gamma(DPPH)$ is the peak-to-peak line-width of the DPPH resonance, and $\Gamma(DPPH )$ is the corresponding line-width of the nitroxide central resonance (Altenbach et al., 1996).

RESULTS

Retention of function following site-specific spin labeling of CaM

Site-directed spin labeling was used to investigate the possible role of the sequence between Met$^{76}$ and Ser$^{81}$ in mediating the structural coupling between the opposing domains of CaM. Five separate CaM mutants were created involving 1) conservative substitutions within the sequence between Met$^{76}$ and Ser$^{81}$ (i.e., T79C or S81C), 2) near the amino terminus of a relatively stable helical structure within the solvent-exposed central sequence between Phe$^{65}$ and Phe$^{92}$ (i.e., F68C, L69C), or 3) within a helical element within the carboxyl-terminal domain in a region expected to undergo significant tertiary contact interactions (i.e., L105C) (Fig. 1). Mutations involving charged amino acids were avoided, as these have previously been shown to stabilize the structure of the central sequence (Sun et al., 1999). In all cases the introduction of individual cysteines and the covalent attachment of MTSSL to these sites results in minimal perturbation of the secondary structure of CaM, because the α-helical content is virtually identical to that of wild-type CaM (Fig. 2; Table 1). Likewise, the CaM dependence of the activation of the plasma membrane Ca-ATPase and the maximal ATPase activity are virtually identical for all five spin-labeled CaM mutants compared with wild-type CaM (Table 1). Thus, the mutation and spin-label modifications do not affect the structure or function of CaM, indicating that spin-label EPR measurements of the rotational dynamics at these positions reflect the native structure of CaM.

Resolution of conformational heterogeneity within the central sequence

In the crystal structure of CaM (1cll.pdb), L69, T79, and S81 do not undergo significant tertiary contact interactions...
with other structural elements and have calculated surface accessibilities of 49, 63, and 52 Å². In contrast, tertiary interactions reduce the solvent accessibilities of Phe68 and accessibility (\(\text{H9016}\)) was measured at 25°C using Eq. 1 in Materials and Methods.

The line-shapes of the spin-label EPR spectra at 25°C are diagnostic of solvent-exposed spin labels, consistent with the solvent-exposed location of these amino acids within the crystal structure of CaM (Babu et al., 1988; Chattopadhyaya et al., 1992; Mchaourab et al., 1996). Addition of 30% sucrose has essentially no effect on the spectral line-shape (data not shown), indicating that the spin-label EPR spectra are insensitive to the rotational motion of the entire CaM molecule and reflect the backbone dynamics of the protein. Decreasing the temperature reduces the rate of motion without significantly affecting the average secondary and tertiary structure of CaM, permitting the molecular anisotropy to be resolved on the spin-label EPR time-scale (Tsalkova and Privalov, 1985; Verheyden et al., 1994; Mchaourab et al., 1996; Sun et al., 2001). Thus, at 4°C the components are clearly separated in the low-field region and indicate the presence of a mobile (\(\beta\)) and motionally restricted (\(\alpha\)) component. The maximal hyperfine splitting of the motionally restricted component is similar at all three positions (i.e., L69C, T79C, and S81C) (Table 1), indicating that the rotational dynamics and environment are similar. However, the fractional contribution of the mobile population of spin labels (\(\beta\)) is larger for T79C and S81C (within the disordered sequence between Met76 and Ser81) relative to L69C within the solvent-exposed central helix that are expected to be within stable helices. Calcium binding enhances the spectral contribution of the motionally restricted component at T79C and S81C, without affecting the maximal hyperfine splitting (Fig. 3 A). Because polarity differences arising from tertiary contact interactions are expected to alter the maximal hyperfine splitting (Mchaourab et al., 1996), these results suggest that the observed spectral changes are sensitive to differences in the secondary structure and backbone dynamics at these solvent-exposed positions.

A consideration of the spin-label EPR spectra at 4°C indicates that irrespective of the spin label position the maximal hyperfine splitting of the EPR spectra is substantially less than 68 G, which is characteristic of a rigid-limit spectrum, suggesting the presence of rotational motion on the nanosecond time-scale (Columbus et al., 2001). This result is consistent with previous fluorescence anisotropy measurements that found fluorophores bound to L69C to undergo restricted rotational motion, with rotational correlation times of 1.9 \(\pm\) 0.3 ns and 13 \(\pm\) 1 ns corresponding to segmental motion and the overall rotational motion of CaM (Sun et al., 1999). Likewise, MTSSL bound to sites within helical elements of T4 lysozyme undergoes restricted rotational motion, with a rotational correlation time of \(\sim\) 2 ns, whose amplitude of motion is affected by the placement of the spin label within the sequence (Columbus et al., 2001). Hubbell and co-workers emphasized that the sensitivity of MTSSL to changes in backbone dynamics is directly related to interactions between the disulfide linkage of the spin label and main-chain atoms within the protein, which restrict the amplitude of motion for the nitroxide moiety (Langren et al., 2001). Thus, within solvent-exposed helices it is expected that MTSSL will selectively monitor backbone fluctuations on the nanosecond time-scale that alters the amplitude of probe motion.

**TABLE 1 Properties of calmodulin mutants**

<table>
<thead>
<tr>
<th>CaM species</th>
<th>ATPase activity ((\mu)mol P(_i) mg(^{-1}) h(^{-1}))</th>
<th>(\alpha)-Helical content</th>
<th>Maximal hyperfine splitting (Gauss)</th>
<th>Line-width ((\Gamma_o)) (Gauss)</th>
<th>(O_2) accessibility (II)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>1.5 (\pm) 0.2</td>
<td>57 (\pm) 1%</td>
<td>apo-CaM ((C^-)^2)CaM</td>
<td>2.1 (\pm) 0.1</td>
<td>0.14 (\pm) 0.2</td>
</tr>
<tr>
<td>F68C</td>
<td>1.3 (\pm) 0.2</td>
<td>57 (\pm) 2%</td>
<td>apo-CaM ((C^-)^2)CaM</td>
<td>2.8 (\pm) 0.1</td>
<td>0.23 (\pm) 0.2</td>
</tr>
<tr>
<td>L69C</td>
<td>1.3 (\pm) 0.1</td>
<td>56 (\pm) 2%</td>
<td>apo-CaM ((C^-)^2)CaM</td>
<td>2.9 (\pm) 0.1</td>
<td>0.17 (\pm) 0.01</td>
</tr>
<tr>
<td>T79C</td>
<td>1.4 (\pm) 0.2</td>
<td>58 (\pm) 2%</td>
<td>apo-CaM ((C^-)^2)CaM</td>
<td>2.4 (\pm) 0.1</td>
<td>0.14 (\pm) 0.01</td>
</tr>
<tr>
<td>S81C</td>
<td>1.5 (\pm) 0.2</td>
<td>59 (\pm) 2%</td>
<td>apo-CaM ((C^-)^2)CaM</td>
<td>2.2 (\pm) 0.1</td>
<td>0.16 (\pm) 0.01</td>
</tr>
<tr>
<td>L105C</td>
<td>1.2 (\pm) 0.1</td>
<td>59 (\pm) 2%</td>
<td>apo-CaM ((C^-)^2)CaM</td>
<td>2.0 (\pm) 0.1</td>
<td>0.11 (\pm) 0.01</td>
</tr>
</tbody>
</table>

Wild-type and CaM mutants were expressed and purified from *E. coli*, as described in Materials and Methods. CaM-dependent rates of ATP hydrolysis for the plasma membrane Ca-ATPase in erythrocyte ghosts was measured in the presence of saturating CaM concentrations, as previously described in detail (Yao et al., 1996). Estimates of \(\alpha\)-helical content were obtained from fits to CD spectra using the program Contin (Venyaminov and Yang, 1996). Maximal hyperfine splitting was measured at 4°C by fitting the spectral extrema to a Gaussian line-shape. Line-width of center peak was measured at 25°C. Oxygen accessibility (II) was measured at 25°C using Eq. 1 in Materials and Methods.
FIGURE 3  Conformational heterogeneity within the central helix is modulated by calcium activation. CaM mutants L69C, T79C, and S81C were individually spin-labeled and spectra were taken at 4°C (A) and 25°C (B) in the apo- and calcium-activated forms. Corresponding spectral features involving the immobilized (α) and mobile (β) spectral components, the maximal hyperfine splitting (arrows), and central field line-width ($\Gamma_o$) are indicated in A and B. For clarity, the high-field extremes are shown expanded 10-fold. Experimental conditions involved 100 μM CaM in 25 mM HEPES (pH 7.5), 0.1 M KCl, 1.0 mM MgCl$_2$, and either 0.1 mM CaCl$_2$ or 0.1 mM EGTA.
The suggestion that MTSSL selectively monitors backbone fluctuations is consistent with the observation that “at the majority of helix surface sites investigated so far, the sequence-specific mobility differences are apparently not determined by differences in interaction with the nearest neighbors in the same helix” (Columbus et al., 2001). Thus, the two component spectra of MTSSL bound to L69C, T79C, and S81C probably represent multiple-protein conformational states. This interpretation is consistent with previous NMR measurements that demonstrated the presence of conformational heterogeneity within the sequence between Met\(^{76}\) and Ser\(^{81}\), which “adopts a helical conformation for about one-third of the time” that is relatively stable on the nanosecond time-scale and may transiently adopt a 3\(_{10}\) conformation (Kuboniwa et al., 1995). Nevertheless, it is significant that two distinct orientations were observed for the disulfide group of MTSSL in the crystal structure of the spin-labeled R119C mutant of T4 lysozyme, suggesting that conformational heterogeneity can arise because of probe heterogeneity (Langren et al., 2000). One conformation corresponds to a relatively mobile component and is suggested to correspond to a conformer in which the disulfide linkage of MTSSL undergoes a S-H non-hydrogen-bonding interaction with Gln at the \(i + 4\) position. This is the only example of the four positions studied in which an interaction with a neighboring side chain resulted in the stabilization of two distinct rotamers of the MTSSL spin label. However, a similar non-hydrogen-bonding interaction between a neighboring Gln in the primary sequence and the disulfide in MTSSL is observed for the K65C mutant, suggesting that interactions between MTSSL and neighboring Gln side chains may result in the appearance of spectral heterogeneity (Langren et al., 2000). The absence of comparable side chains in the vicinity of the sequence between Met\(^{76}\) and Ser\(^{81}\) makes it unlikely that corresponding interactions occur between MTSSL and proximal side chains in CaM. Nevertheless, it remains possible that the two component spectra of MTSSL bound to sites within CaM could result from probe heterogeneity that is unrelated to protein structural heterogeneity.

Trifluoroethanol stabilizes the motionally restricted component

To investigate the possible relationship between the two motional populations observed in the spin-label EPR spectra and protein secondary structural elements, we have investigated the dynamic structure of the S81C spin-labeled CaM mutant (which exhibits a substantial mobile component) in the presence of trifluoroethanol (TFE), which has previously been shown to stabilize \(\alpha\)-helical structures in a range of proteins including CaM (Bayley and Martin, 1992; Evans, 1995; Luo and Baldwin, 1997). In fact, the use of nonaqueous solvents that stabilize \(\alpha\)-helical structures has been suggested to result in the stabilization of the helix between Met\(^{76}\) and Ser\(^{81}\) in the crystal structure of CaM (Babu et al., 1988; Bayley and Martin, 1992).

We find that the addition of as little as 3% (v/v) TFE stabilizes the motionally restricted conformation for both apo- and calcium-activated CaM (Fig. 4). No alteration in the maximal hyperfine splitting occurs, indicating that at these concentrations TFE selectively stabilizes the immobilized component without significantly affecting the polar environment in the vicinity of the spin label. In contrast, substantially larger amounts of TFE are required to induce line-shape changes in the EPR spectra of MTSSL at F68C and L105C (Fig. 5), which are within more stable helical regions of CaM (Kuboniwa et al., 1995; Zhang et al., 1995; Finn et al., 1995). These results demonstrate a differential sensitivity of MTSSL bound within the sequence between Met\(^{76}\) and Ser\(^{81}\) relative to more stable secondary structural

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**FIGURE 4** Calcium and TFE stabilize the central helix. Spectra for the apo- and calcium-activated forms of the spin-labeled CaM mutant S81C are shown in the absence and presence of 3% (v/v) TFE. High-field extremes are shown expanded 10-fold. Experimental conditions involved 100 \(\mu\)M CaM in 25 mM HEPES (pH 7.5), 0.1 M KCl, 1.0 mM MgCl\(_2\), and either 0.1 mM CaCl\(_2\) or 0.1 mM EGTA. Temperature was 4°C.
elements within either the amino- or carboxyl-terminal domains of CaM. Because denaturation of CaM following the addition of guanidine hydrochloride enhances the fractional contribution of the mobile component (β) (data not shown), these results further suggest that there is a dynamic equilibrium within this sequence between a mobile conformation and a motionally restricted conformation that are in slow exchange on the spin-label EPR time-scale. Thus, in the context of earlier NMR results that suggest the sequence between Met76 and Ser81 adopts a helical conformation approximately one-third of the time (Kuboniwa et al., 1995), these results suggest that the immobilized component reflects the helical conformation within this sequence. Because calcium binding stabilizes the immobilized component, these results further suggest that calcium binding mediates a direct coupling between the opposing domains of CaM through the stabilization of the helical content between Met76 and Ser81.

**Tertiary contact interactions modify maximal hyperfine splitting and solvent accessibility**

In comparison with MTTSL bound to L69C, T79C, and S81C, calcium-dependent changes in the maximal hyperfine splitting
Splitting are evident for spin-labeled CaM mutants F68C and L105C (Fig. 5). These results are consistent with the presence of tertiary contact interactions between side chains at these positions within the crystal structure of CaM (Babu et al., 1988; Chattopdhyaya et al., 1992) and suggest that L69C represents the first residue within the central sequence that does not undergo significant tertiary contact interactions with side chains in other secondary structural elements. Before calcium binding, the rotational mobility of the mobile spectral component ($\beta$) for spin labels covalently bound to positions 68 and 105 is larger relative to those bound at solvent-exposed positions (i.e., L69C, T79C, and S81C), as is evident from their larger spectral intensity and the smaller spectral line-widths associated with the center field resonance ($I^{c}_{0}$) (Figs. 3 and 5; Table 1). The enhanced rotational dynamics of the mobile component at these sites may reflect the ability of MTSSL to access a variety of conformations that minimize steric interactions with other side chains at buried sites (Langren et al., 2000). Alternatively, the enhanced mobility may reflect main-chain segmental movements that reflect the partial unwinding of the $\alpha$-helices at positions 68 and 105, which are, respectively, located at the amino terminus of helices within the crystal structure of CaM (Babu et al., 1988; Chattopdhyaya et al., 1992). Consistent with this latter hypothesis, the addition of 15% (v/v) TFE enhances the spectral contribution of the immobilized fraction of spin labels (Fig. 5A). Likewise, the CaM sequence near Leu$^{105}$ has previously been demonstrated to form a noncompact and disordered state with a highly susceptible proteolytic cleavage site at Arg$^{106}$ (Newton et al., 1984; Tsalkova and Privalov, 1985; Kuboniwa et al., 1995; Shea et al., 1996). These latter results suggest that to be accessible to the active site of the protease that the backbone fold exhibits some nonhelical behavior.

**DISCUSSION**

**Structure and function of the central sequence**

The structural coupling between the opposing domains of CaM was originally evident in the crystal structure of CaM. This structure demonstrates two homologous globular domains that include amino acid residues that are part of a central sequence, which exists as an extended $\alpha$-helix between Phe$^{65}$ and Phe$^{92}$ (Babu et al., 1985, 1988; Chattopdhyaya et al., 1992) (Fig. 1). However, the existence of a stable central sequence in the cell remains controversial, because inclusion of nonaqueous solvents during crystallization stabilizes the helical content within the central sequence (Ikura et al., 1991; Barbato et al., 1992; Bayley and Martin, 1992). Furthermore, under cellular conditions, techniques that monitor the solution structure have demonstrated static disorder within the central sequence that results in a distribution of conformations that modifies the spatial arrangement of the opposing domains (Heidorn and Trewhella, 1988; Yao et al., 1994; Tjandra et al., 1995; Kuboniwa et al., 1995). It is, therefore, often argued that the central sequence functions as a flexible tether, whose major function is to maintain the spatial proximity between the opposing domains necessary for target protein binding (Persechini and Kretsinger, 1988; Kretsinger, 1992; Tjandra et al., 1999). Despite the flexibility of the central sequence, at physiologically relevant ionic strength and pH the central sequence assumes conformationally distinct structures in the apo-form and following calcium activation (Small and Anderson, 1988; Yao et al., 1994; Sorensen and Shea, 1996; Wrighers et al., 1998; Sun et al., 1999). In this respect, mutations that affect the structure of the central sequence diminish the binding affinity between CaM and target proteins (Craig et al., 1987; Kataoka et al., 1991; Sacks et al., 1996; Yin et al., 2000b; Sun et al., 2001). These latter results suggest that stabilization of the central sequence may function to minimize nonspecific interactions between the opposing domains of CaM involved in binding to target proteins. The calcium-dependent stabilization of the structure of the central sequence between Met$^{76}$ and Ser$^{81}$ observed in the present study, therefore, have important implications with respect to the mechanism of target protein binding by CaM.

**Conformational heterogeneity within the central sequence of CaM**

In this study we have used site-directed spin labeling coupled with the introduction of unique cysteines to resolve calcium-induced conformational changes within the sequence between Met$^{76}$ and Ser$^{81}$ in vertebrate CaM. Two component spectra were observed for MTSSL spin labels covalently bound at positions T79C and S81C in the solvent-exposed interdomain central sequence of CaM using spin-label EPR spectroscopy (Fig. 3), which is indicative of heterogeneity involving the backbone dynamics of CaM. Calcium activation, which was reproduced by the addition of as little as 3% (v/v) TFE, enhances the spectral contribution associated with the immobilized component at positions T79C and S81C (Figs. 3 and 4). Because TFE has previously been demonstrated to stabilize the $\alpha$-helical content of CaM and other proteins (Bayley and Martin, 1992; Evans, 1995; Luo and Baldwin, 1997), the immobilized component probably reflects a population of helical secondary structures (within the interdomain sequence between Met$^{76}$ and Ser$^{81}$) that are stabilized by calcium binding. In contrast, much larger amounts of TFE are required to induce line-shape changes at positions F68C or L105C, consistent with their greater conformational stability (Fig. 5). These results are consistent with previous suggestions that the sequence between Met$^{76}$ and Ser$^{81}$ is “delicately balanced between helical and non-helical conformations” (Kuboniwa et al., 1995) and adopts multiple conformations in which a relatively long-lived helical conformation is assumed for

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about one-third of the time in apo-CaM (Kataoka et al., 1991; Kretsinger, 1992; Kuboniwa et al., 1995; Wriggers et al., 1998).

The underlying reasons for conformational heterogeneity within the central sequence have been suggested to involve both 1) the transient disruption of specific hydrogen bonds, which can induce the unwinding of helical elements, and 2) a reorientation of the opposing globular domains that favors target protein binding (Wriggers et al., 1998). The presence of multiple conformations within the central sequence that undergo slow exchange on the nanosecond time-scale is, furthermore, consistent with previous suggestions that conformational rearrangements between the opposing domains of CaM are slow relative to the time-scale of overall protein rotational motion, which has been measured using both fluorescence anisotropy and NMR spectroscopy to be between 9 and 12 ns at 20°C (Small and Anderson, 1988; Török et al., 1992; Barbato et al., 1992; Yao et al., 1994; Sun et al., 1999). Thus, in the context of these prior observations, the current results demonstrating that a significant fraction of the interdomain central sequence is conformationally restricted provides strong evidence that the calcium-dependent structural linkage between the opposing domains of CaM involves alterations in the structure of the interdomain central sequence between Met76 and Ser81.

Significance of central sequence to high-affinity CaM binding

Previous measurements have demonstrated that initial binding between CaM and target proteins normally involves the high-affinity association of the carboxyl-terminal domain of CaM followed by association of the amino-terminal domain. Association of the amino-terminal domain is aided by the reduced volume available for its diffusion following the main. Association of the amino-terminal domain is facilitated by minimizing interdomain interactions. Consistent with this suggestion, a reduced binding affinity to target proteins is apparent in CaM variants with sequence differences within calcium-binding site four that results in the loss of high-affinity calcium binding and structural interactions between the opposing domains (Lee and Klevit, 2000; Yin et al., 2000b). Likewise, the binding affinity between CaM and target proteins is diminished following the structural uncoupling between the opposing domains by oxidative modification of selected methionines near the carboxy terminus or the elimination of the hydrogen bond between Trp138 and Glu82 (Gao et al., 1998; Yin et al., 2000a; Sun et al., 2001). All of these results support earlier suggestions that the interdomain sequence is marginally stable under normal cellular conditions (Kretsinger, 1992) and that calcium binding to the carboxyl-terminal domain stabilizes the helical content of the central sequence. Binding of the carboxyl-terminal domain to target proteins appears to function as a conformational switch that is required for binding of the amino-terminal domain. Thus, following target protein association, stabilizing interactions between sites within the carboxyl-terminal domain and the central sequence are disrupted, resulting in the independent rotational dynamics of the amino-terminal domain that is necessary for target protein association and the structural collapse of CaM around the CaM-binding sequence.

Conclusions and future directions

We have demonstrated calcium-dependent structural changes involving the stabilization of a more rigid structure within the interdomain sequence involving Met76 to Ser81. To clarify the proposed role of stabilizing interactions between the carboxyl-terminal domain and the central sequence in mediating the high-affinity interaction between CaM and target proteins, future measurements should identify how alterations in the interdomain sequence affects the kinetics of CaM binding to target proteins.

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