Structure and Dynamics of Calmodulin in Solution

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ABSTRACT  To characterize the dynamic behavior of calmodulin in solution, we have carried out molecular dynamics (MD) simulations of the Ca²⁺-loaded structure. The crystal structure of calmodulin was placed in a solvent sphere of radius 44 Å, and 6 Cl⁻ and 22 Na⁺ ions were included to neutralize the system and to model a 150 mM salt concentration. The total number of atoms was 32,867. During the 3-ns simulation, the structure exhibits large conformational changes on the nanosecond time scale. The central α-helix, which has been shown to unwind locally upon binding of calmodulin to target proteins, bends and unwinds near residue Arg⁷⁴. We interpret this result as a preparative step in the more extensive structural transition observed in the “flexible linker” region 74–82 of the central helix upon complex formation. The major structural change is a reorientation of the two Ca²⁺-binding domains with respect to each other and a rearrangement of α-helices in the N-terminus domain that makes the hydrophobic target peptide binding site more accessible. This structural rearrangement brings the domains to a more favorable position for target binding, poised to achieve the orientation observed in the complex of calmodulin with myosin light-chain kinase. Analysis of solvent structure reveals an inhomogeneity in the mobility of water in the vicinity of the protein, which is attributable to the hydrophobic effect exerted by calmodulin’s binding sites for target peptides.

INTRODUCTION

The calcium-sensing protein calmodulin (CaM) exemplifies the function of protein domain movements in the regulation of cellular processes. CaM is a small protein (Babu et al., 1988; Chattopadhya et al., 1992) consisting of 148 amino acid residues that belongs to a class of ubiquitous proteins with similar structure characterized by their distinctive helix-loop-helix Ca²⁺-binding motif, the so-called EF hand (Kretsinger, 1980). CaM is involved in the regulation of most of the important Ca²⁺-dependent signaling pathways in the cell (Cohen and Klee, 1988; James et al., 1995). Fig. 1 shows that in the crystal, CaM adopts a dumbbell conformation in which the long helix connecting the two Ca²⁺-binding domains is completely extended. In this conformation the two domains are in a trans-like orientation. Early small-angle x-ray scattering (SAXS) results suggested that the tethering helix in CaM is flexible, allowing the two domains to adopt various relative orientations and separations in solution (Seaton et al., 1985; Heidorn and Trewhella, 1988; Matsushima et al., 1989). These results motivated computational studies of both CaM and the related protein troponin C that also supported a flexible interdomain linker (Mehler et al., 1991). The molecular dynamics (MD) simulations suggested that large regions of conformational space were accessible to CaM, allowing it to adopt a continuum of relative separations and orientations, ranging from the fully extended structure observed in the crystal to a very compacted structure suggested by the SAXS studies when CaM was complexed to small peptides like mellitin or mastoparan (Kataoka et al., 1989; Matsushima et al., 1989). This flexibility was also supported by early NMR results (Barbato et al., 1992). The functional significance of this ability of CaM to achieve a large range of relative conformations of its N-terminal and C-terminal domains became clear from the crystal structure of CaM complexed with a peptide analog of smooth muscle myosin light-chain kinase (Meador et al., 1992) and by the solution structure of calmodulin with a peptide analog of skeletal muscle myosin light-chain kinase (Ikura et al., 1992). In both of these CaM-peptide complexes a short helical segment at the interface of helices D and E (Fig. 1) was found to unwind, leading to a compacted protein with reoriented domains in cis-like conformation, where the hydrophobic patches face each other and hold the peptide between them (reviewed by Weinstein and Mehler, 1994).

Recently, NMR structures of Ca²⁺-free (apo) calmodulin were reported (Finn et al., 1995; Zhang et al., 1995; Kuboniwa et al., 1995), which, in comparison with the crystal structures of Ca²⁺-saturated (holo) calmodulin (Babu et al., 1988; Chattopadhya et al., 1992), revealed the details of the conformational transition induced by Ca²⁺ binding to the protein. In both domains of holo calmodulin, in which all four Ca²⁺-binding sites are occupied, a large, hydrophobic surface is exposed to the solvent (Tanaka and Hidaka, 1980). These hydrophobic patches, which are nearly completely buried in apo-CaM, form the binding interface with α-helices of CaM’s target proteins and contribute most of the binding energy (Ikura et al., 1992; Meador et al., 1992). Each hydrophobic surface is surrounded by a rim of mostly negatively charged residues, which may act to orient the target sequences by interactions with positively charged
residues of the target proteins and contribute to the binding energy in the complex (Zhang et al., 1995).

The early simulations of CaM in solution suggested two types of structural changes relative to the crystal structure: a reorientation of the domains and an overall compaction of the protein (Weinstein and Mehler, 1994). Both changes have been observed in complexes of CaM with targets, as pointed out above, but it was not clear if both are required to prepare isolated CaM for the interaction with a target peptide. The incomplete solvent model used in the earlier simulations (Pascual-Ahuir et al., 1991) may have led to artifactual structural changes not involved in preparing CaM for target interaction. In this paper we seek the molecular basis of the experimentally well-characterized structural flexibility of CaM, as reviewed above, in the context of a full description of the protein-solvent system. With the help of the simulation of the molecular dynamics of CaM, it becomes possible to exhibit the molecular mechanisms that control the structural flexibility and its relation to function, including the role of solvent. Furthermore, analysis of the molecular rearrangement of CaM from the structure in the crystal to that in solution offers insights about the subtle intrinsic properties that determine preparation for target binding, as discussed recently (Weinstein and Mehler, 1994, 1996).

Results of the simulation reveal how the protein-solvent interactions act to modulate the properties of the protein and of the aqueous ionic solution. These offer an indication of the dependence of CaM function on both explicit and bulk interactions with the solvent. In ionic solution, calmodulin’s negative charge (−15e) is balanced by counterions. Most of the Na+ counterions can be expected to form a diffuse atmosphere of ions surrounding the protein (Israelachvili, 1992). We determined the distribution of ions as they explore the accessible conformational space in the MD simulations and compared the results to a continuum model of ion distribution derived from a numerical solution of the Poisson-Boltzmann equation (Honig and Nicholls, 1995). The dynamics of the solvent emerging from the analysis of the long trajectory identifies a change in the mobility of water molecules near the surface of CaM. Although protein-water interactions have been the subject of intense computational research (Teeter, 1991; Daggett and Levitt, 1993), it is still unclear how exposed hydrophobic side chains alter the dynamics of the surrounding water. Recently, MD simulations suggested a reduced number of hydrogen bonds in the first solvation shell of hydrophobic compounds (Laidig and Daggett, 1996), confirming Muller’s “modified hydration-shell hydrogen-bond” model for the molecular explanation of the hydrophobic effect (Muller, 1988, 1990). This model is opposed to the classic “clathrate cage” model of water hydrogen bonds (Powell, 1948), which predicts increased order near hydrophobic surfaces relative to bulk solvent. Of special interest, in this respect, are the hydrophobic target binding patches in the N-terminal and C-terminal domain of CaM (Ikura et al., 1992). In this paper we report deviations of translational water diffusion near calmodulin’s hydrophobic patches and compare the results with other theoretical and computational studies of protein-solvent interactions.
COMPUTATIONAL METHODS

Preparation of structures, solvent model, and molecular dynamics

The coordinates of Ca$^{2+}$-loaded calmodulin were obtained from PDB entry 3CLN (Babu et al., 1988). For the simulation, residues 1–4 and 148, which are missing in the crystal structure, were added as described previously (Pascual-Ahuir et al., 1991). The completed protein was then immersed in a sphere of explicit water molecules with a radius of 44 Å. The water shell was constructed by covering the system with a three-dimensional lattice of water cubes provided by X-PLOR (Brünger, 1992). To neutralize the system and to model physiological ionic strength, 22 sodium and 6 chloride ions were iteratively placed at the minima and maxima of the electrostatic potential surrounding the protein, following the method of Mark et al. (1991). This results in a system with a nominal ionic strength of 150 mM, comprising 32,867 atoms (10,474 water molecules), shown in Fig. 1.

To prepare the system the solvent was first equilibrated with fixed protein coordinates. After energy minimization, initial velocities were assigned according to a Maxwell distribution, and the solvent was heated to 310 K in steps of 30 K in a 5-ps time period, and then equilibrated for 60 ps at 310 K. The total energy of the final system was minimized, followed by heating to 310 K and equilibration for 40 ps. Free MD simulation was carried out for 3 ns. The parameters of the PAR19 united atom force field of CHARMM (Brooks et al., 1983) were used for the protein with default values of the control parameters, unit dielectric constant, and a 1-fs integration step. Ion parameters were obtained from the PAR22 force field for Na$^+$ and Cl$^-$ and from Hori et al. (1988) for Ca$^{2+}$. The TIP3(P) water model (Jorgensen et al., 1983) was modified by relaxing internal geometry constraints to provide water flexibility. This modification was motivated by MD studies that demonstrated an improvement of physical properties of flexible water over the rigid TIP3 model (Teeter, 1991; Daggett and Levitt, 1993). Flexible water models such as F3P (Levitt and Sharon, 1988) and F3C (Levitt, 1989) better reproduce dynamical water properties compared to TIP3, which overestimates the translational diffusion coefficient by up to 60% (Daggett and Levitt, 1993; Obst and Bradaczek, 1996). Flexible TIP3P water exhibits values for the water density and heat of vaporization as close to the experimental values as those obtained with the original TIP3P model (Steinbach and Brooks, 1993). Furthermore, the structure of water and the ionic hydration shells is little influenced by the flexibility of the TIP3 water molecules (Guàrdia and Padró, 1996).

Structural and statistical analysis

Fluctuations

Because the two globular regions are joined by a flexible linker that permits a high degree of relative motion of the domains as two rigid bodies, it is necessary to remove this motion to compare the calculated B-factors and order parameters with the experimental results. Two least-squares fits (Kabsch, 1976) were carried out for each entry in the trajectory to remove the rigid body motions of the N-terminal (res. 1–77) and C-terminal (res. 78–148) domains, respectively. The atomic fluctuations calculated from the simulation were compared with the B-factors in the crystal from the PDB entry 3CLN (Babu et al., 1988), using the relationship

$$ \langle \Delta r_i^2 \rangle = 3B_i / (8\pi^2), $$

where $\Delta r_i$ is the atomic displacement of atom $i$ from its average position and $B_i$ is the corresponding B-factor. The fluctuations reported are averages of the backbone atom fluctuations for each residue, as described below.

Conformational variability

The conformational changes of the protein during the simulation were described by the root mean square (rms) deviations from chosen reference structures. A linear measure is the time-dependent rms value that gives the departure of the protein from the starting, crystal-like structure. To monitor the time evolution of the simulation structure and its convergence in phase space, mutual rms deviations (rsm$\alpha$) were calculated for coordinate sets $\bar{r}(t)$ of either the N- or the C-terminal domain, using the relationship

$$ \text{rms}_\alpha = \sqrt{ \frac{1}{N_\alpha} \sum_{i=1}^{N_\alpha} [r_{\alpha}(t_i) - \bar{r}_{\alpha}(t_j)]^2 }, $$

where $\bar{r}_{\alpha}(t_i)$ and $\bar{r}_{\alpha}(t_j)$ are the least-squares fitted (Kabsch, 1976) coordinate sets of $\alpha$-carbons ($\alpha = (1, 2, \cdots, N_\alpha)$) at times $t_i$ and $t_j$, and where $N_\alpha$ is the number of residues in the domain of interest (77 and 71 for the N- and C-terminal domains, respectively).

$^{15}$N order parameters

The orientational procedure described above results in atomic coordinates that are expressed in domain-fixed reference systems. This procedure corresponds to the model assumptions made by Barbato et al. (1992) to determine the order parameters from their NMR measurements. Thus the values calculated here can be directly compared with the experimental results. The $^{15}$N order parameters were calculated according to the model free formulation of Lipari and Szabo (1982).

Identifying rigid domains

Hinge-bending movements, where rigid domains are connected by flexible joints, are the dominating type of conformational changes documented in the Brookhaven Protein Data Bank (Gerstein et al., 1994). By means of the new Hingefind algorithm (Wriggers and Schulten, 1997), we
examined the movements of calmodulin’s structural domains that can be considered to undergo local noise. The HingeFIND algorithm compares two known conformations of a protein and identifies connected regions that exhibit preserved packing within a specified tolerance of positional fluctuations. These relatively rigid regions are found in an iterative procedure, in which poor matching residues are excluded from a selected subset of a-carbons and good matches are included. After dividing the protein into rigid domains, the algorithm determines effective rotation axes (hinges) that characterize the movements of the identified domains relative to each other. The identification of hinge axes and of corresponding rotation angles provides a reduced representation of the complex movements exhibited by the protein. We extracted rigid domains by comparing calmodulin’s structure after a 3-ns simulation to the initial crystal structure. A 2.0-Å tolerance in the position of a-carbons was chosen to filter, as described (Wriggers and Schulten, 1997), the significant movements from local noise and thermal disorder.

Counterion distribution

Localized clusters in the distribution of sodium counterions were determined from a three-dimensional histogram. Trajectory frames from the production run were least-squares fitted (Kabsch, 1976) to the initial protein structure every 100 fs. The positions of the 22 sodium ions in each of the fitted frames were recorded in bins of (1.4 Å)³ size. The resulting density map was normalized such that the sum over all histogram densities equals the total number of sodium ions.

The electrostatic potential around CaM was calculated with GRASP (Nicholls et al., 1991). A dielectric constant of 2 was assigned to the protein interior, and 80 was used for water. The charges were taken from the PAR19 parameter set of CHARMM (Brooks et al., 1983), resulting in a net charge of −15e on the protein including the calcium ions. The linear Poisson-Boltzmann equation (McQuarrie, 1976; Honig and Nicholls, 1995) was solved at various ionic strengths, and the concentration of sodium counterions was determined from the electrostatic potential calculated from the Boltzmann factor at temperature T = 310 K.

Diffusion coefficients

Translational diffusion coefficients can be calculated from the mean square displacement function

\[ 6D = \langle [\mathbf{r}(\tau + t) - \mathbf{r}(\tau)]^2 \rangle + C, \]  

where \( \langle \cdot \cdot \cdot \rangle \) denotes an average over time origins \( \tau \) and over a species of molecules of interest, \( \mathbf{r}(t) \) is the position of a molecule at the offset time \( t \), and \( C \) is a constant accounting for short-time effects in the dynamics of investigated molecules (Wong et al., 1989; Sansom et al., 1996; Breed et al., 1996). For short offset times the diffusive model of motion breaks down: at the picosecond time scale the distance traveled by a molecule is only on the order of interatomic distances, and atom displacements may originate to a significant degree from local fluctuations in the evolving system. Hence, in simulations with periodic boundary conditions, \( D \) is usually estimated from the long time limit of Eq. 3 (Eriksson et al., 1995; Obst and Bradaczek, 1996). In our case of confined diffusion, we evaluated \( D \) at short offset times \( t \) to prevent finite system size effects. Time offsets 10 ps < \( t \) < 100 ps were found to be practical to avoid both finite size effects and short-time correlations. We obtained mean values and standard deviations of the diffusion coefficients by evaluating Eq. 3 for this time window.

Water mobility

The three-dimensional distribution of water mobility was evaluated as follows. Trajectory frames from the production run were least-squares fitted (Kabsch, 1976) to the initial protein structure every picosecond. The displacements of water molecules were recorded in (3.0 Å)³ histogram bins. The traveling water molecules force a trade-off between the feasible spatial resolution of the diffusion coefficient and the possible length of the offset time \( t \). Hence displacements were only recorded for short offset times \( t_1 = 1 \) ps and \( t_2 = 2 \) ps. The short-time mobility of water surrounding the protein was characterized by the spatial distribution of displacements for \( t = t_1 \). To compare the short-time mobility with the diffusion coefficient determined by Eq. 3, an effective diffusion coefficient \( D_{\text{eff}}^{(1)} \) was computed from

\[ 6D_{\text{eff}}^{(1)}(i, j, k) = \left\langle [\mathbf{r}(\tau + t_i) - \mathbf{r}(\tau)]^2 \right\rangle_{ijk}, \]

where \( \langle \cdot \cdot \cdot \rangle_{ijk} \) denotes an average over time origins \( \tau \) and over a histogram bin at point \( r_{ijk} \) on the three-dimensional grid of width 3.0 Å.

The diffusive contribution to the local motion of water molecules, \( D_{\text{eff}}^{(2)} \), was estimated from the slope of the mean square displacement function, averaged over the recorded displacements in the histogram bins at offset times \( t_1 = 1 \) ps and \( t_2 = 2 \) ps (Lounnas et al., 1994),

\[ 6D_{\text{eff}}^{(2)}(i, j, k)(t_2 - t_1) = \left\langle [\mathbf{r}(\tau + t_2) - \mathbf{r}(\tau)]^2 \right\rangle_{ijk} - \left\langle [\mathbf{r}(\tau + t_1) - \mathbf{r}(\tau)]^2 \right\rangle_{ijk}, \]

where \( \langle \cdot \cdot \cdot \rangle_{ijk} \) denotes an average over time origins \( \tau \) and over a histogram bin at point \( r_{ijk} \) on the three-dimensional grid of width 3.0 Å.

RESULTS AND DISCUSSION

Dynamics and stability

In the course of the 3-ns MD simulation, CaM undergoes several conformational changes relative to the initial crystal structure that appear to be completed after \( \sim 1.5 \) ns (Fig. 2). During the second half of the simulation, the system fluctuates around a new average position at 5.2 (± 0.6) Å rms
deviation from the initial structure. Although the overall rms deviation steadily increases in the first 1.5 ns, the rms deviations of the individual domains remain much smaller (Fig. 2). The structural fluctuations of the C-terminal domain (res. 78–147) converge rapidly within 100 ps, and the structure remains close to the crystal structure (1.3 (± 0.1) Å rms deviation). The N-terminal domain (residues 5–77) also reaches a plateau after about ~100 ps (1.5 (± 0.1) Å rms deviation), where it remains for ~1300 ps. Subsequently, the system undergoes a conformational change that is characterized by a rapid change in rms deviation between 1.3 and 1.5 ns, and then stabilizes after ~2 ns at 2.3 (± 0.1) Å rms deviation (Fig. 2).

A second type of structural rearrangement is the reorientation of the two domains from the crystal structure to a configuration closer to that observed in CaM-target complexes. This is seen in Fig. 3, where the virtual dihedral angle (VDA) between the four calcium ions is plotted against the simulation time. The VDA was shown earlier to be a reasonable representation of the interdomain orientation (Pascual-Ahuir et al., 1991). In contrast to the conformational change in the N-terminal domain, the reorientation of the two Ca-binding domains relative to each other is a gradual process that occurs during the first 1.5 ns of the simulation. The VDA starts at the value measured in the crystal structure (~134°) and then steadily decreases to an average value around ~200° (160° in the standard IUPAC convention). This latter value of the VDA is much closer to that assumed by CaM when it is complexed to targets (~110° in the IUPAC convention; Ikura et al., 1992). More-
over, comparison of the temporal behavior of the VDA with the global rms deviation plotted in Fig. 2 shows that the VDA stabilizes at about the same time as the rms deviation, which suggests that the relative orientation of the two domains underlies the time dependence of the global rms deviation.

The time evolution of the structural changes can be further characterized by using a two-dimensional representation of the mutual rms deviations of trajectory frames (Eq. 2). Fig. 4 reveals distinct features in the sampling of the configurational space associated with each globular domain. The trajectory of the C-terminal domain corresponds to a single family of structures with mutual rms deviations of < 1.3 Å. The more flexible N-terminal domain exhibits several stages of conformational changes. During a first stage that lasts for ~1.4 ns, the N-terminal domain remains close to the initial conformation. Subsequently, after increasing fluctuations in the structure starting at 900 ps simulation time, the N-terminal domain undergoes an abrupt conformational change. After 1.4 ns, the system continues to exhibit large fluctuations until ~2 ns simulation time, and subsequently appears to settle into the new conformation for the remainder of the 3-ns trajectory. As Fig. 4 shows, the mutual rms deviations of the N-terminal domain structures during the last nanosecond of the simulation do not exceed 1.3 (± 0.2) Å, where the standard deviation is the maximum value computed for each structural region.

The evolution of the conformational changes as shown in Fig. 4 suggests that the trajectory can be divided into a relaxation phase (0–2 ns) and a quiescent phase (2–3 ns). In the relaxation phase the protein responds to the absence of the intermolecular interactions present in the crystal lattice (Chattopadhyaya et al., 1992), and then the system fluctuates around a new average structure without undergoing further structural rearrangements during the remainder of the trajectory (Figs. 2 and 3). Because the C-terminal domain is also involved in a large number of intermolecular interactions in the crystal structure (Chattopadhyaya et al., 1992), an eventual change of conformation cannot be ruled out, especially in view of the functional significance of the structural change observed in the N-terminal domain (see below). From Fig. 4 it is seen that the structure of the C-terminal domain late in the trajectory is moving further away from the structure early in the trajectory. The increasing difference among the structures after ~2.8 ns is reminiscent of the similar increase noted in the N-terminal domain after ~900 ps that preceded the structural change.
Chain flexibility

Atomic fluctuations were sampled over three time intervals: 0–1 ns, 1–2 ns, and 2–3 ns (Fig. 5). The patterns of internal fluctuations observed in the three intervals are fairly similar for most regions of the protein, indicating the convergence of this property. One exception was the region comprising residues 39–61 that exhibit increased mobility during the second nanosecond of the simulation. These increased fluctuations probably reflect the conformational change exhibited by the N-terminal domain at 1.5 ns (Fig. 2), and the large fluctuations in the N-terminal domain that preceded and followed the structural change (Fig. 4). Two regions, comprising residues 37–42 and 110–115, exhibit high mobilities in the simulation relative to the rest of the protein. This enhanced local flexibility may be due to missing contacts in the crystal lattice, e.g., of residues 37–42 of protein (000) with residues 114–120 of protein (011), defined by Chattopadhyaya et al. (1992). Moreover, these two six-residue segments lie in homologous regions in the N-terminal and C-terminal domains (Strynadka and James, 1988), suggesting a possible functional significance of the observed increased fluctuations.

The computed fluctuations are compared with those estimated from the crystal B-factors (Babu et al., 1988) in Fig. 5. As was also found for other molecular systems (McCammon and Harvey, 1987), there is a good qualitative agreement between the fluctuations calculated from the last nanosecond of the MD run and the experimental values. On the average, and especially in the C-terminal domain, the fluctuations observed from the crystal structure are larger than the values calculated from the simulation, which is probably due to a certain static disorder in the crystal unrelated to intrinsic movements (Kuriyan and Weis, 1991). Because of the flexible connection of the two globular domains, they behave as quasi-independent bodies (Barbato et al., 1992), and therefore the static disorder in each of them can be quite different.

Order parameters

Global conformational changes may complicate identification of the intrinsic motional freedom of the protein backbone from the computed fluctuations. Another descriptor of the absolute backbone motion is obtained from the relaxation of bond rotation. Here the autocorrelation functions and generalized order parameters were calculated for each backbone amide bond. The autocorrelation function is given by (Lipari and Szabo, 1982)

\[ C_\alpha(t) = \langle P_2 (\mathbf{v}_\alpha(\tau) \cdot \mathbf{v}_\alpha(t + \tau)) \rangle, \]  

where \( P_2 \) is the second-order Legendre polynomial, \( \mathbf{v}_\alpha(t) \) is the instantaneous N-H unit vector of residue \( \alpha \) in the protein fixed-frame reference system, and the angle brackets indicate time averaging. In the “model free” approach proposed by Lipari and Szabo (1982), the relaxation described by \( C_\alpha(t) \) is encapsulated by two parameters: a generalized order parameter and the relaxation time. The order parameter, \( S^2 \), is a measure of the restriction of the angular motion of the N-H bond vector. Its value ranges from 1, for completely hindered motion, to 0 for completely free, isotropic motion. The order parameter of each N-H bond \( \alpha \) is calculated from the asymptotic limit of its time autocorrelation function (Lipari and Szabo, 1982), i.e.,

\[ S^2 = \lim_{t \to \infty} C_\alpha(t) = \frac{4\pi}{5} \sum_{m=-2}^{2} |Y_{2m}(\Omega)|^2, \]  

where \( Y_{2m} \) are the second-order spherical harmonics and \( \Omega \) is the orientation of the N-H bonds relative to a macromolecular fixed frame.
Autocorrelation functions and order parameters were calculated for all N-H bond vectors from the final nanosecond of the simulation, because during this interval no major changes in the structural properties occur. Similar to results from previous studies of proteins (Chandrasekhar et al., 1992; Eriksson et al., 1993), most residues were found to undergo fast motions that are efficiently sampled in the course of the simulation. The extended length of the present trajectory made it possible to monitor slow relaxation events for 28 residues, as evidenced by the slow convergence of \(C_n(t)\). Only four residues were found to undergo motions that are insufficiently sampled on the nanosecond time scale. The validity of the simulation was further assessed by comparing the computed order parameters with experimental NMR results for CaM (Barbato et al., 1992).

The \(C_n(t)\) of most of the N-H bond vectors showed a sharp initial decay, followed by a plateau that extends over the rest of the 460-ps sampling window used. This indicates that these N-H groups undergo small-amplitude, fast librational motions along well-defined directions. The 28 residues exhibiting slower relaxation events that converged on the 300-ps time-scale are mostly located in flexible parts of the protein and do not participate in intramolecular interactions. Residues 70–82 of the tether belong to this group of slowly convergent \(C_n(t)\) and were found to undergo large structural fluctuations that mediate the relative motion of the two globular domains. The corresponding values of the order parameters of residues 77–81 are significantly larger than those observed experimentally. Finally, the correlation functions of residues Glu\(^3\), Gly\(^61\), Thr\(^26\), and Glu\(^45\) did not converge.

The N-H order parameters have been averaged over secondary structural elements; they are given in Table 1. Overall, the trends observed from the measured values are fairly well reproduced; the \(\alpha\)-helical order parameters exhibit the strongest restriction, whereas the intradomain linkers show the highest degree of rotational freedom. The average errors between measured and calculated order parameters for the N-terminal and C-terminal domains are 0.006 and 0.07, respectively, and the rms differences are 0.02 and 0.07, respectively, so that the calculated order parameters for the N-terminal domain are in slightly better agreement with experiment than those for the C-terminal domain. Comparison of the measured order parameters (Barbato et al., 1992) in the two domains shows that, on average, the C-terminal domain has a slightly greater rotational flexibility than the N-terminal domain, in agreement with the main-chain B-factors (Babu et al., 1988). This difference between the two domains was not resolved by the simulation. It is finally noted that the calculation does not reproduce the experimentally observed low order parameters of residues 57 and 130 located at homologous positions of loop 2 and loop 4, respectively. Instead, the calcium-binding loops exhibit uniformly highly restricted motions.

### Domain movements

The structural origins of the conformational changes predicted by the simulation were investigated by comparing the structure after a 3-ns simulation against the initial structure in terms of rigid domains identified with the algorithm Hingefind, as described in Computational Methods. To determine the domains, the structure after 3 ns was averaged over the last 20 ps of simulation time and then energy-minimized. Fig. 6 presents a backbone trace of this structure compared to the initial structure.

At 2.0-Å resolution, the C-terminal lobe comprises the largest rigid domain found, whereas the N-terminal lobe splits into four rigid subdomains. The two structures were superimposed by their rigid C-terminal domains, and all subdomain translations and rotations were evaluated relative to this reference orientation. The effective rotation axes determined with Hingefind permit a coarse representation of the exhibited conformational changes. Helix A (brown in Fig. 6) reorients and moves toward the C-terminal domain. The binding site of calcium 1 with helix B (blue), as well as helix C (yellow), rotates about axes perpendicular to the central helix, thereby opening up and extending the structure. An inspection of the “animated” protein dynamics revealed that these movements are prominent in the first nanosecond of the simulation.

The major conformational change in the N-terminal domain that occurs around 1.5 ns (Fig. 2) causes the calcium 2 binding site (green) to swing to the side by a 65° rotation about the central helix (Fig. 6). This unwinding of the helix avoids steric clashes with the central helix when other rigid parts (calcium 1 binding site with helix B, and helix C) move to open up the N-terminal domain (Fig. 6). It appears that conformational strain in the structure also produces a kink in the central alpha helix at Met\(^{12}\).

### Table 1: Average order parameters for secondary structural elements of CaM

<table>
<thead>
<tr>
<th>Element</th>
<th>Residues</th>
<th>Simulation</th>
<th>NMR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Helix A</td>
<td>5–19</td>
<td>0.84</td>
<td>0.84</td>
</tr>
<tr>
<td>Helix B</td>
<td>29–38</td>
<td>0.86</td>
<td>0.83</td>
</tr>
<tr>
<td>Helix C</td>
<td>45–55</td>
<td>0.85</td>
<td>0.83</td>
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<tr>
<td>Helix D</td>
<td>65–76</td>
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<td>0.85</td>
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<tr>
<td>Helix E</td>
<td>82–92</td>
<td>0.86</td>
<td>0.78</td>
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<tr>
<td>Helix F</td>
<td>102–111</td>
<td>0.87</td>
<td>0.82</td>
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<td>Helix G</td>
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<td>Helix H</td>
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<td>Linker 1</td>
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<td>0.75</td>
</tr>
<tr>
<td>Linker 2</td>
<td>112–117</td>
<td>0.75</td>
<td>0.64</td>
</tr>
</tbody>
</table>

Comparison of simulation and NMR results (Barbato et al., 1992). The average values do not include those residues for which the correlation functions did not converge during the simulation, and for which there are no experimental data.
The effect of the conformational changes in the N-terminal domain on the solvent exposure of the hydrophobic target peptide binding patch is shown in Fig. 7. The reorientation as described by the change in VDA (Fig. 3) amounts to a movement of the N-terminal domain from a trans-like orientation in the crystal structure toward a cis-like conformation required for target binding. Although the positioning of the reactive surface of the N-terminal domain of the simulated structure at 3 ns (Fig. 7) suggests that this surface may be more accessible to target peptides, its solvent-accessible surface area (SASA), calculated using a standard probe of radius 1.4 Å, is actually less than the SASA from the crystal structure. This is true of the C-terminal reactive surface as well. However, the radius associated with a target peptide will be considerably larger than that of a water molecule, and the surface area accessible to a probe representing such a ligand may be considerably different from the SASA. The dependence of the accessible surface area (ASA) on probe size was evaluated, therefore, and is shown in Fig. 8 for both domains. Comparison of the crystal structure shows a decreasing ASA for the domains as the probe radius is increased. This type of behavior of the ASA, which also applies to the structure at 1 ns, i.e., before the structural change in the N-terminal domain has occurred, indicates that the reactive surfaces are concave. However, at 2 ns, after the structural change, the response of the ASA to probe size is different in the two domains: the ASA of the N-terminal reactive surface (top panel in Fig. 7) increases strongly with increasing probe size, whereas in the C-terminal (bottom panel in Fig. 7) it continues to show behavior similar to that at 1 ns and to that of the crystal structure. The increase in ASA of the N-terminal domain is even larger at 3 ns, but here again, the ASA in the C-terminal surface remains unchanged. The change in response of the ASA in the N-terminal domain indicates that the reactive surface has changed from concave to convex, with probe radii of 10–15 Å experiencing increases in ASA by a factor of 2 to 3 relative to the crystal structure.

The ability of the N-terminal domain of CaM to relax to conformations better suited to interaction with targets sug-
gests a functional role of its greater flexibility. A more pronounced flexibility of the N-terminal domain relative to the C-terminal domain was also observed, albeit to a lesser degree, in calculations on the isolated domains (Mehler and Weinstein, unpublished results). In contrast, the C-terminal domain remains relatively rigid (Figs. 2 and 5) during the 3-ns simulation, but as mentioned above, the C-domain may eventually undergo conformational changes as well, to bring its reactive surface into a more favorable configuration for interaction with target peptides.

Conformational changes in the tether

In addition to the observed kink at Met72, a further structural change associated with the reorientation of the N-terminal domain is an unwinding of the central α-helix at residue Arg74. This is indicated by the replacement of the typical \( i, i + 4 \) α-helical hydrogen bonding pattern of the crystal structure with an \( i, i + 5 \) hydrogen bond between Arg74 and Leu89. The disruption of the helix was first observed after 900 ps of simulation time, a little more than halfway through the reorientation process (Fig. 3). Fig. 9 demonstrates that the unwinding of the α-helix is associated with a perturbation of torsion angles in residues 71–73; in particular, the \( \Psi \) torsional angles in residues 71 and 72 show large deviations from the accepted range. An unwinding of the helix by one residue yields a relative rotation of 72° of the two separated parts of the α-helix. This angular change is close to the observed change in the VDA (Fig. 3) and the rotation of the calcium 2 binding site relative to the C-terminal domain (Fig. 6). The unwinding of the flexible...
linker at Arg$^{74}$ observed in the simulation appears to facilitate the reorientation of the two calcium-binding domains of CaM.

In earlier simulations (Mehler et al., 1991; Pascual-Ahuir et al., 1991; Weinstein and Mehler, 1992) three arginine residues (Arg$^{74}$, Arg$^{86}$, and Arg$^{90}$) were found to form and sever H-bonds with successive attachment sites from more distant parts of the structure, leading to compaction and interdomain reorientation. Of these three arginines, Arg$^{74}$ was most clearly involved in the structural reorganization that included both reorientation of the domains and overall compaction of the system. In contrast, the present simulation shows no compaction of the calmodulin structure: despite the conformational changes, the radius of gyration of the protein remained stable at a value of 22 Å. The results indicate a functional role for Arg$^{74}$ in the bending and unwinding of the flexible linker. In the simple solvent models used in the earlier studies, the region between the two domains was free of water and thus exhibited a low dielectric volume. Consequently, it is likely that in the earlier simulations the structure responded in an artifactual way that was similar to the bending response elicited by the presence of target peptide, because the latter will remove water from the region between the domains (Elcock and McCammon, 1996). Nevertheless, it is noteworthy that simulations on the isolated tethering helix indicate that it is inherently flexible (Mehler et al., 1991; van der Spoel et al., 1996), and therefore it remains possible that simulations substantially longer than the current 3 ns may lead to further conformational changes in the flexible linker that would result in structural changes in the protein in the absence of target peptides.

Solvent properties

Water makes up 86% of the total mass of the simulated system. The solvent was modeled in the absence of boundary conditions (Roberts and Schnitker, 1995), and long-range electrostatic forces were truncated at 8.5 Å. These restrictions, necessary to achieve a reasonable speed of calculation, need to be applied with care (Roberts and Schnitker, 1995; Del Buono et al., 1994). We show below that the dynamic solvent properties agree well with results from generally accepted benchmark simulations and corroborate the validity of the water force field and solvent modeling approach employed in our simulation. The analysis also reveals a remarkable variability of water mobility that originates in functionally significant protein-solvent interactions.

Counterion distribution

The trajectory permits an improved sampling of the counterion distribution surrounding the protein. Earlier simula-
tions of biopolymer-water systems have suffered from short simulation times that allowed the sampling of only a small fraction of the total accessible ion configuration space (Eriksson et al., 1995). To characterize this sampling, we compare the time-averaged distribution of sodium ions calculated from the 3-ns trajectory with the ion concentration obtained from solution of the Poisson-Boltzmann equation (McQuarrie, 1976; Honig and Nicholls, 1995). Fig. 10 presents the three-dimensional histogram of sodium ions sampled from the trajectory and the local ion concentration $\rho_{Na}$ calculated from the electrostatic potential surrounding the protein, namely

$$\rho_{Na}(r) = \rho_o \exp \left( -\frac{\Xi(\rho_o, \vec{r})}{k_B T} \right), \quad (8)$$

where $\rho_o$ is the ionic strength and $\Xi(\rho_o, \vec{r})$ is the potential. The figure demonstrates that the counterions are well distributed and more localized in regions of negative electrostatic potential. We estimated the effective bulk ionic strength $\rho_o$ of the system to be 75 mM, which is half of the total ion concentration ([Na] + [Cl]) in our model. An inspection of computed distributions $\rho_{Na}(\vec{r})$ for values of the ionic strength ranging from 10 mM to 150 mM showed that the local density is insensitive to changes in the bulk ionic strength $\rho_o$.

The local densities calculated from the simulation are larger and the distribution is more scattered than predicted by the continuum model for the diffuse counterion cloud (Fig. 10). The truncation of long-range electrostatic interactions may have favored short-range ion-ion and ion-protein interactions as the ions localized in regions of high sodium density near the protein surface, and undersampling may have caused the observed sodium density fluctuations; it is possible, albeit not certain, that longer simulation times than the current 3 ns may produce a better agreement with the continuum model.

Diffusion of solvent molecules

The statistical properties of aqueous ionic solutions, such as solvent-solvent and solvent-ion pair distribution functions, are insensitive to changes in boundary conditions (Roberts

FIGURE 10 Counterion condensation observed in simulation and predicted by electrostatic theory. Calmodulin is represented by its solvent-accessible surface averaged from a 2–3-ns simulation time. (A) Three-dimensional histogram of sodium ion distribution in calmodulin’s vicinity sampled from the 3-ns trajectory using a (1.4 Å)$^3$ bin size. The colors code for ion densities corresponding to a concentration $\rho_{Na} = 1.0$ M (red), and $\rho_{Na} = 3.0$ M (yellow). (B) Contours of the sodium concentration $\rho_{Na}(\vec{r})$ and the electrostatic potential $\Xi(\rho_o, \vec{r})$ (Eq. 8) near the surface of the negatively charged (-15e) protein. The contours were calculated with the program GRASP (Nicholls et al., 1991) at ionic strength $\rho_o = 0.075$ M. The following contours are shown: $\rho_{Na} = 0.1$ M, $\Xi = -0.29 k_B T$ (blue), $\rho_{Na} = 0.2$ M, $\Xi = -0.98 k_B T$ (green), and $\rho_{Na} = 0.5$ M, $\Xi = -1.90 k_B T$ (yellow).
and Schnitker, 1995), the water model (Guàrdia and Padró, 1996), and the model of Coulomb interactions (Del Buono et al., 1994), in the sense that the calculated properties all are close to the experimental values. In contrast, these and other authors (Sansom et al., 1996; Teeter, 1991; Daggett and Levitt, 1993) have shown that dynamical solvent properties, such as translational diffusion coefficients, are model dependent and may deviate significantly from the experimental values. To test the reliability of our solvent model and of the force field parameterization, we calculated the diffusion coefficients of water molecules, sodium ions, and chloride ions. In Fig. 11 the mean square displacement functions of these solvent molecules are compared. The resulting values of the diffusion coefficients, as determined from Eq. 3, were $D_{\text{water}} = 2.3 \pm 0.1 \times 10^{-9}$ m$^2$/s, $D_{\text{Na}} = 0.55 \pm 0.03 \times 10^{-9}$ m$^2$/s, and $D_{\text{Cl}} = 1.1 \pm 0.1 \times 10^{-9}$ m$^2$/s. The experimental value of pure water at 300 K is $D_{\text{water}} = 2.3 \times 10^{-9}$ m$^2$/s (Eisenberg and Kauzmann, 1969) and can be extrapolated to $D_{\text{water}} = 2.9 \times 10^{-9}$ m$^2$/s at the simulation temperature of 310 K (Obst and Bradaczek, 1996). The translational mobility of water molecules obtained above for our protein-solvent system is slightly smaller than the experimentally measured value. The agreement with the experimental value is good compared to simulations of pure TIP3 water ($D_{\text{water}} = 4.0 \times 10^{-9}$ m$^2$/s; Jorgensen, 1981). The observed sodium ion diffusion is low, possibly because of the preference of sodium ions for positions near the negatively charged protein surface (Fig. 10). The value of $D_{\text{Na}}$ obtained here is close to the value $0.5 \times 10^{-9}$ m$^2$/s for simulated Na$^+$ near DNA (Eriksson et al., 1995). (As with water, one would expect a slight increase in ionic diffusion at 310 K relative to 300 K. Here we discuss only experimental and computational values obtained for the more common temperature 300 K.) Other simulations resulted in a value of $D_{\text{Na}} = 1.2-1.3 \times 10^{-9}$ m$^2$/s (Norberg and Nilsson, 1994; Obst and Bradaczek, 1996), and the experimental datum is $D_{\text{Na}} = 1.33 \times 10^{-9}$ m$^2$/s (Hertz, 1973). The calculated diffusion of the chloride ions is also low compared to the experimental value of $2.03 \times 10^{-9}$ m$^2$/s (Tyrell and Harris, 1984). Values of $D_{\text{Cl}} = 0.5-2.2 \times 10^{-9}$ m$^2$/s (Roberts and Schnitker, 1995; Guardia and Padrò, 1996) have been reported for simulated Cl$^-$ diffusion. Our estimate of $D_{\text{Cl}}$ near the protein, therefore, appears reasonable.

The diffusion coefficient $D$, as determined by Eq. 3, has a physical meaning only for homogeneous and isotropic systems, in which the solvent mobility is independent of the positions visited by each solvent molecule. Consequently, for an inhomogeneous system such as a solvated protein, Eq. 3 only describes an average property of the system. A physically more meaningful representation of the solvent mobility in the vicinity of a protein should take into account the spatial dependence of the diffusive motion. As described in Computational Methods, it is possible to compute the mobility of solvent molecules in localized bins of a three-dimensional histogram, if the distance traveled within the sampling time $t$ (Eq. 3) is small compared to the bin size used.

Fig. 12 presents contours of constant rms displacements of water molecules for $t = 1.0$ ps. The short offset time allows one to determine the distribution of the depicted displacements at a resolution smaller than the 3.0-Å bin size used to calculate the global value of $D$. Water-protein interactions and the water model used cause a spatial inhomogeneity of observed water mobility. An increase in mobility near the surface of the solvent sphere is artifactual, because of the absence of bulk solvent that would surround the spherical system. A decrease in mobility to values as low as $D_{\text{eff}}^{(1)} = 2.0 \times 10^{-9}$ m$^2$/s can be attributed to the coordination of water molecules by the polar surface of the protein. An inspection of the distribution of $D_{\text{eff}}^{(1)}$ values reveals that the hydrophobic effect exerted on the solvent

![FIGURE 11 Mean square displacements, $\langle (\vec{r}(t + \tau) - \vec{r}(t))^2 \rangle$, of water molecules (solid line), sodium ions (dashed line), and chloride ions (dotted line) calculated from the 3-ns trajectory. The displacement functions allow one to compute corresponding diffusion coefficients (Eq. 3).](image)
induces an enhanced mobility to values as high as $D_{\text{eff}}^{(1)} = 10^{\alpha} \cdot 10^{-9}$ m$^2$/s at calmodulin’s hydrophobic target-peptide binding patches.

The observed short-time water mobility corresponds to an effective diffusion coefficient $D_{\text{eff}}^{(1)}$ (Eq. 4) that encompasses both diffusive and nondiffusive contributions to the short-time water motion. The average of $D_{\text{eff}}^{(1)}$ over all histogram bins (Fig. 12) yields a water diffusion coefficient $D_{\text{water}} = 2.9 \cdot 10^{-9}$ m$^2$/s, which is in closer agreement with the value of $2.3 \cdot 10^{-9}$ m$^2$/s determined above. Therefore $D_{\text{eff}}^{(2)}$ represents a good approximation of the diffusive contribution to the water motion.

**SUMMARY AND CONCLUSION**

The present MD simulation of Ca$^{2+}$-saturated calmodulin in solution has revealed several structural features relevant to calmodulin’s ability to form complexes with target proteins. Relaxation from the crystal constraints allows the central helix to bend and unwind at residue Arg$^74$. The simulation yields a water diffusion coefficient $D_{\text{water}} = 2.9 \cdot 10^{-9}$ m$^2$/s, which is in closer agreement with the value of $2.3 \cdot 10^{-9}$ m$^2$/s determined above. Therefore $D_{\text{eff}}^{(2)}$ represents a good approximation of the diffusive contribution to the water motion.

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conformational change in the protein actin (Chik et al., 1996). A slight difference in energy between the open and the collapsed state revealed an extreme sensitivity of the actin conformation to changes in the chemical and thermal environment of the crystal (Chik et al., 1996).

After relaxation from the crystal constraints, the calmodulin structure stabilized during the latter part of the simulation trajectory. In this quiescent phase of the simulation, the structure fluctuated about a new average conformation, although it is unlikely that the conformational relaxation from the crystal was completed. As discussed recently, sampling errors due to short simulation times preclude the identification of slow modes in the dynamics of proteins (Balsera et al., 1996). Likewise, in the 3-ns simulation, we do not expect to have sampled all possible movements of calmodulin. The two individual domains, however, should be well equilibrated, because of the small system size and the relatively long simulation time.

The relatively long simulation times in the fully solvated system that are required for the appearance of the structural changes associated with CaM function demonstrate that this process cannot be characterized with short simulations of a few hundred picoseconds. Rather, trajectories of at least several nanoseconds are required, and even these cannot provide a full account of the details in the molecular dynamics that control the relationship between structure and function in CaM. These findings are likely to be generalizable to other systems that undergo large structural changes related to their function.

Although the structure-function relationship of calmodulin has been the subject of intense research (Weinstein and Mehler, 1994), one basic question was not answered fully. This question concerns the mechanism whereby calmodulin is able to bind and activate a variety of distinct proteins in the cell. A considerable body of evidence indicates that calmodulin can assume multiple modes of interaction with targets other than the modes realized in the complexes with myosin light chain kinase (Ikura et al., 1992; Meador et al., 1992). For example, in the γ-subunit of phosphorylase kinase, two noncontiguous peptides (Phk5 and Phk13) form complexes with calmodulin. Whereas Phk5 has an α-helical conformation in the complex, it appears that Phk13 forms an elongated complex, in which the peptide is bent into a hairpin-shaped structure (Juminaga et al., 1993). The spacing of hydrophobic residues of cyclic nucleotide phosphodiesterase (Charbonneau et al., 1991) and calmodulin kinase II (Bennet and Kennedy, 1987) suggests that their binding to calmodulin is different.

Some of the intrinsic properties of calmodulin that enable it to realize multiple target binding modes have been iden-
tified. Experimental observations indicate an asymmetry in target affinity of calmodulin’s domains. The tryptic fragment corresponding to the C-terminal domain binds preferentially to target peptides (Newton et al., 1984, 1985), at affinities more than 100 times higher (Sanyal et al., 1988), compared to the fragment corresponding to the N-terminal domain. Our results reveal that calmodulin’s C-terminal domain is relatively rigid, whereas the central α-helix and the N-terminal domain are flexible. This relative ordering in the rigidity of the two domains supports a binding mechanism in which the C-terminal domain binds to its target first, followed by the N-terminal domain (Persechini et al., 1994). The inherent flexibility of both the N-terminal domain and the tethering linker allows a suitable binding conformation once the second binding site is recognized. The nature and size of the hydrophobic patches involved in target binding depend on this flexible reorientation and constitute elements of recognition specificity.

A large number of computational and experimental studies focused on the behavior of water molecules near the surface of solvated proteins (Teeter, 1991; Daggett and Levitt, 1993). Such studies indicate that the dynamic properties of water close to the protein are perturbed, but there is some disagreement on whether water molecules are immobilized (Wong and McCammon, 1986; Brooks and Karplus, 1989; Levitt and Sharon, 1988; Tirado-Rives and Jorgensen, 1990) or hypermobile (Lounnas et al., 1994) relative to bulk water. Our results indicate a uniformly reduced diffusion coefficient of water near calmodulin. However, in terms of short-time mobility, we need to differentiate between water molecules near the polar surface of the protein and water molecules near the hydrophobic patches. Our results demonstrate an increased short-time mobility of waters near the hydrophobic patches, rooted in nondiffusive motion. Other authors have reported an effect of hydrophobicity on the diffusive motion relative to polar residues, but there is disagreement on whether hydrophobic surfaces increase (Levitt and Sharon, 1988) or decrease (Karplus and Rossky, 1980; Brooks and Karplus, 1989) the diffusion coefficient. Clearly, the disagreement between the computational studies suggests that the simulation of the physical properties of water at a protein surface are model-dependent because of limitations of the various protein and water force fields.

For realistic simulations of solvated proteins, molecular dynamics force fields require refinements that will yield better agreement with the experimentally observed behavior of aqueous solutions. Of special interest for such refinements are water molecules near the protein surface, which have been elucidated by recent protein crystal structures. It was hypothesized (Klotz, 1958) that water near hydrophobic residues would form clathrate cages (Powell, 1948), but a study of water distributions in 16 protein structures with resolutions below 1.7 Å did not reveal such patterns (Thanki et al., 1988). A more recent model of the hydrophobic effect suggests a reduced number of hydrogen bonds in the first solvation shell of hydrophobic compounds (Muller, 1988, 1990). This model has been supported by computer simulations (Laidig and Daggett, 1996). Water molecules with a reduced number of hydrogen bonding partners should be kinetically more labile. In this respect Muller’s hypothesis is consistent with the enhanced mobility of water molecules for short times and small displacements observed in this work for calmodulin’s hydrophobic patches. On the other hand, the classical picture of clathrate cages seems to pertain to the average structure of the hydrogen-bonding network, which is not inconsistent with the existence of kinetically labile waters at a hydrophobic surface.

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