Backbone Dynamics of a Symmetric Calmodulin Dimer in Complex with the Calmodulin-Binding Domain of the Basic-Helix-Loop-Helix Transcription Factor SEF2-1/E2-2: A Highly Dynamic Complex

Göran Larsson,* Jürgen Schleucher,* Jacqueline Onions,† Stefan Hermann,† Thomas Grundström,† and Sybren S. Wijmenga*

*Department of Medical Biochemistry and Biophysics, and †Department of Molecular Biology, University of Umeå, S-901 87 Umeå, Sweden

ABSTRACT Calmodulin (CaM) interacts specifically as a dimer with some dimeric basic-Helix-Loop-Helix (bHLH) transcription factors via a novel high affinity binding mode. Here we report a study of the backbone dynamics by 15N-spin relaxation on the CaM dimer in complex with a dimeric peptide that mimics the CaM binding region of the bHLH transcription factor SEF2-1. The relaxation data were measured at multiple magnetic fields, and analyzed in a model-free manner using in-house written software designed to detect nanosecond internal motion. Besides picosecond motions, all residues also experience internal motion with an effective correlation time of ~2.5 ns with squared order parameter (S0) of ~0.75. Hydrodynamic calculations suggest that this can be attributed to motions of the N- and C-terminal domains of the CaM dimer in the complex. Moreover, residues with significant exchange broadening are found. They are clustered in the CaM:SEF2-1mp binding interface, the CaM:CaM dimer interface, and in the flexible helix connecting the CaM N- and C-terminal domains, and have similar exchange times (~50 μs), suggesting a cooperative mechanism probably caused by protein:protein interactions. The dynamic features presented here support the conclusion that the conformationally heterogeneous bHLH mimicking peptide trapped inside the CaM dimer exchanges between different binding sites on both nanosecond and microsecond timescales. Nature has thus found a way to specifically recognize a relatively ill-fitting target. This novel mode of target-specific binding, which neither belongs to lock-and-key nor induced-fit binding, is characterized by dimerization and continuous exchange between multiple flexible binding alternatives.

INTRODUCTION

Calmodulin (CaM) is a Ca2+ binding protein, present in all eukaryotic cells. It has a 100% amino acid identity among all analyzed vertebrates, and plays a central role in translating intracellular Ca2+ signals into biological responses.

The crystal structures of Ca2+/CaM show an extended dumbbell shaped molecule, in which its two globular domains are connected with a long α-helix (1). In solution, this helix is disrupted in the middle (2), which allows N- and C-terminal domains of both apo-CaM and Ca2+/CaM to tumble almost independently of each other (3–5). Upon Ca2+ binding, a conformational change enables it to interact with over 100 different target proteins, including transcription factors (for reviews, see Van Eldik and Watterson and others (6,7)).

When Ca2+/CaM interact with its targets, the flexibly connected domains normally collapse into a more compact globule. This is called "wraparound" binding because the two domains in CaM wrap around the α-helical target (8,9). In context of the wraparound binding mode, Wand and co-workers have studied the backbone and side-chain dynamics of free CaM and CaM in complex with a peptide (10). Upon peptide binding, the flexibility of side chains located in the binding sites is redistributed. This is believed to assist the target-specific deformation of the binding sites in CaM that is necessary for productive binding. In contrast, the backbone within the domains is fairly rigid, both for target-free (3) and target-bound CaM (10). Tjandra and co-workers (3) have shown, via 15N-relaxation at multiple fields, that in free CaM the two domains undergo a slow "wobbling" motion on a timescale of ~3 ns, apparently not evident in target-bound CaM.

Although the wraparound mode of target binding is the most studied, alternative binding modes have become evident. One of these, the interaction of CaM with basic-Helix-Loop-Helix (bHLH) transcription factors has a 2:2 stochiometry (11). This new type of CaM interaction is the first example where two interacting CaM molecules interact with a dimeric target.

Helix-Loop-Helix (HLH) transcription factors regulate numerous developmental processes (12,13). Most HLH proteins belong to the bHLH group, which has a basic sequence directly N-terminal to the HLH motif. They are active as dimers, where the two basic regions bind DNA symmetrically as α-helices on opposite sides in the major groove.
In the absence of DNA, the basic sequences lose their well-defined secondary structure (14).

Ca\(^{2+}\) signaling can inhibit the transcriptional activities in vivo of the bHLH proteins E12 and SEF2-1 through direct binding of Ca\(^{2+}/\)CaM to the basic region of the proteins, resulting in inhibition of their DNA binding (15). Thus, CaM can, in a Ca\(^{2+}\)-dependent manner, directly interact with some members of the bHLH family. The CaM binding site coincides with the basic DNA-binding sequence of the bHLH dimers (11,15,16,17). A homodimeric peptide corresponding to the complete SEF2-1 basic sequence, henceforth called SEF2-1mp, was chosen as a good model system for NMR studies of the interactions between bHLH proteins and Ca\(^{2+}/\)CaM.

In an earlier NMR study we could conclude that the CaM:SEF2-1mp complex has a 2:2 stoichiometry, where two interacting CaM molecules bind one homodimeric SEF2-1mp (11). The previous NMR data were consistent with two schematic models of the CaM dimer. In both models the N-terminal domain of one CaM faces the C-terminal domain of the other CaM, creating a hydrophobic tunnel where SEF2-1mp is trapped. Fig. 1 shows one of these two possible models. In the same study we could also conclude that SEF2-1mp lacks any well-defined secondary structure when interacting with CaM, an observation very unusual for CaM-bound peptides. The peptide still interacts with the same exposed hydrophobic patches as in the wraparound binding mode, but here a number of weak interactions occur instead of one strong. Despite these features the overall interaction is highly specific with nanomolar binding strength (16). Possibly the dimeric nature of both the target and CaM overcomes the less specific hydrophobic interactions. Study of the dynamics of the complex can give insights into this novel type of CaM:target interaction.

In this report we present the backbone dynamics of the CaM dimer bound to the dimeric SEF2-1mp. The CaM:SEF2-1mp complex is found to be highly flexible with internal motions on the picosecond, nanosecond, and microsecond timescales. The dynamics is similar to that found for the “wraparound” binding mode in some aspects, e.g., rigid domains, but also uniquely different in other aspects. The interaction neither belongs to the category of lock-and-key nor to that of induced-fit.

**MATERIALS AND METHODS**

**Sample preparation**

SEF2-1mp mimics the DNA- and CaM-binding region of the bHLH transcription factor SEF2-1. It is a homodimeric peptide formed via a disulfide bridge between two cysteine residues located at position 19 of the 21-residue-long peptide strands. Azodicarboxylic acid (diamide) was used to oxidize the cysteines to form the disulfide bridge and was present in all NMR samples. That the peptide remained dimeric was also confirmed by SDS-PAGE. The preparation of the CaM:SEF2-1mp NMR samples has been described by us earlier (11).

**NMR spectroscopy**

If nothing else is mentioned, the NMR measurements were performed at 308 K. The experiments were carried out on Bruker DRX-400, AMX2-500, and DRX-600 spectrometers equipped with triple-resonance \(\text{H}^1/\text{C}^{13}/\text{N}^1\) or broadband) probes with XYZ-gradient capabilities. The spectra were processed with XWINNMR (Bruker Instruments, Billerica, MA). Proton chemical shifts were calibrated using the internal standard DSS (0.0 ppm at 308 K). \(\text{N}^1\) chemical shifts were indirectly referenced using the gyromagnetic ratio of \(\text{N}^1/\text{H}^1\) (18).

Longitudinal \(R_1\) and transverse \(R_2\) \(\text{N}^1\)-spin relaxation rates were measured at 400 and 600 MHz \(\text{H}^1\) frequency (40.5 and 60.8 MHz \(\text{N}^1\) frequency) using standard pulse sequences (19). The \(\text{N}^1\) \(R_1\) and \(R_2\) experiments at each field were recorded in an interleaved manner with the relaxation delays randomly distributed. The experimental details of these experiments are compiled in Supplementary Materials (Table RS1). Before the start of each experiment the temperature was calibrated with a water/DSS sample.

The \(\text{N}^1\)–(\(\text{H}^1\)) nuclear Overhauser enhancement (NOE) shows only weak field dependence and was therefore only carried out at the higher field. NOE values were determined from pairs of spectra recorded interleaved with and without a 4-s proton saturation (see Supplementary Materials, Table RS1, for more details).

To determine the presence of conformational exchange on microsecond and millisecond timescales, the relaxation-compensated CPMG experiment (20) was carried out at 308 K at 500 MHz and at 308 and 300 K at 600 MHz. The delay between the 180° pulses (\(\delta\)) in the CPMG was set to either 450 \(\mu\)s or 3.6 ms. More experimental details are found in Supplementary Materials (Table RS1).

**Data analysis**

All spectra were analyzed using SYBYL software (TRIPOS). The relaxation data at 600 MHz were integrated over an elliptically shaped area with diameters 7.2 and 6.6 Hz in the \(\text{H}^1\) and \(\text{N}^1\) dimension, respectively. The 400-MHz data were analyzed with \(\text{H}^1\) and \(\text{N}^1\) diameters of 6.5 and 4.4 Hz. The use of small integration areas has the advantage that the noise is still averaged, while at the same time partially overlapping crosspeaks still can be reliably integrated (21).

\(R_1\) and \(R_2\) values were determined by fitting peak volumes to a two-parameter single exponential decay using MATLAB. The error in the \(R_1\) and...
The equations for the relaxation of a backbone amide $^{15}$N spin in a protein as well as their interpretation in terms of the model-free approach of Lipari and Szabo (25) are well described in the literature (see, e.g., Clore et al. and others (23,26,27,28,29)). For the analysis of the relaxation data, the apparent overall tumbling time is an important parameter and we briefly describe the relevant equations.

The apparent overall correlation time at a certain magnetic field $B$, $(\tau_{\text{m}}^0)_{\text{app}}$, is calculated from the ratio $R_2$ over $R_1$ at that field (22,29):

$$ (\tau_{\text{m}}^0)_{\text{app}} = \frac{1}{a_{\text{N}}} \left( \frac{3}{2(1 + a)} \left( \frac{R_2^0}{R_1^0} - \frac{7}{6} (1 + a) \right) \right), \quad (1) $$

with $a = -0.02$. In absence of internal motion and exchange broadening, we define $(\tau_{\text{m}}^0)_{\text{app}}$ as $(\tau_{\text{m}}^0)_{\text{corr}}$ (22). For isotropic tumbling, $(\tau_{\text{m}}^0)_{\text{app}}$ is equal to the true uniform overall correlation time $\tau_{\text{m}}^0$. For anisotropic tumbling, $(\tau_{\text{m}}^0)_{\text{app}}$ contains information on the orientation of the N-H$^\text{N}$ relaxation vector in the molecular frame, which can be used to determine the diffusion tensor (22,26,30,31). For small degrees of anisotropy and axial symmetry the orientation information in $(\tau_{\text{m}}^0)_{\text{app}}$ depends only on the angle, $\Phi$, of the N-H$^\text{N}$ relaxation vector relative to the long axis of the diffusion tensor (22) according to:

$$ (\tau_{\text{m}}^0)_{\text{app}} = \frac{\tau_{\text{m}}^0}{1 + \frac{\Delta}{2} \sin^2(\Phi)}, \quad (2) $$

Here the $\Delta$ is anisotropy and is given by $\Delta = \tau_{\text{m}}^0/\tau_{\text{m}}^0 - 1$, where $\tau_{\text{m}}^0$ is the tumbling time of the long axis, and $\tau_{\text{m}}^0$ is the tumbling time of the short axis of the molecule.

In presence of internal motion, $(\tau_{\text{m}}^0)_{\text{app}}$ depends on the magnetic field, timescale, and degree of internal motion (22,32). We have developed a method for analyzing $^{15}$N-spin relaxation measured at two magnetic fields that corrects for internal motion up to $\sim 4$ ns, giving $(\tau_{\text{m}}^0)_{\text{app-corr}}$ (22). In the absence of internal motions slower than $\sim 4$ ns, $(\tau_{\text{m}}^0)_{\text{app-corr}}$ can thus be considered equal to $(\tau_{\text{m}}^0)_{\text{app}}$, the real overall tumbling time. In the presence of even slower internal motions it becomes progressively difficult to distinguish overall tumbling and internal motion (22,32). Consequently, $(\tau_{\text{m}}^0)_{\text{app-corr}}$ may contain contributions from these slow internal motions and must be considered as an effective overall tumbling time.

For studies of conformational exchange it is important to ascertain whether the exchange is in the fast or slow exchange limit. Usually, it is possible to distinguish between these limits by the number of resonances per exchanged spin, present in the NMR spectra. It is to be noted that observation of single resonances in NMR spectra does not necessarily mean that the exchange is fast. This problem has been considered and recipes on how to still estimate the timescale of an exchange process based on CPMG data can be found in the literature (26,33,34). Below we consider equations assuming fast exchange. In addition, using the simple general equation, which covers for both fast and slow exchange, derived by Ishima and Torchia (34), we also show that the exchange rate can be determined from a combination of CPMG measured $R_{\text{ex}}$ and change in $R_{\text{ex}}$, ($\Delta R_{\text{ex}}$, vide infra) without prior assumption on the exchange timescale.

For a nucleus exchanging between two states, A and B, with different chemical shifts, the apparent exchange broadening, $R_{\text{ex}}$, measured using a CPMG sequence (26) is given by:

$$ R_{\text{ex}} = \frac{\rho_A \rho_B}{\omega_I} \frac{\Delta^2}{\omega_I^2} \left( 1 - \frac{\tan(h(k_{\text{ex}} \delta))}{k_{\text{ex}} \delta} \right). \quad (3) $$

Here, $\rho_A$ and $\rho_B$ are the populations of the states A and B, $\Delta_{\text{ex}} = \Omega_A - \Omega_B$ is the chemical shift difference between the two states, $k_{\text{ex}} = k_{A \rightarrow B} / p_B = k_{B \rightarrow A} / p_A$ is the rate constant for the exchange process, $\delta$ is the delay of the $\delta$-180° CPMG block and $\omega_I$ is the frequency of nucleus I. With the method of Loria et al. (20), $R_2$ is measured for two settings of $\delta$ (we used $2 \delta_1 = 450 \mu$s and $2 \delta_2 = 3.6$ ms) and $R_{\text{ex}}$ is detected if the difference in $R_2$ between the two $\delta$ settings, $\Delta R_{\text{ex}}$, is significantly large.

$$ \Delta R_{\text{ex}} = R_2(\delta_2) - R_2(\delta_1) = \frac{\rho_A \rho_B \Delta^2}{k_{\text{ex}}} \frac{\omega_I}{\tan(h(k_{\text{ex}} \delta_2)) - \tan(h(k_{\text{ex}} \delta_1))}. \quad (4) $$

As can be seen from Eq. 5, the ratio of $\Delta R_{\text{ex}}$ and $R_{\text{ex}}$ depends only on $k_{\text{ex}}$. 

Biophysical Journal 89(2) 1214–1226

Larsson et al.
\[ \frac{\Delta R_{ex}}{R_{ex}} = \left( \frac{\tan(h(k_{ex}\delta_2) - \tan(h(k_{ex}\delta_1))}{k_{ex}\delta_2} - \frac{\tan(h(k_{ex}\delta_1))}{k_{ex}\delta_1} \right) \left( 1 - \frac{\tan(h(k_{ex}\delta_1))}{k_{ex}\delta_1} \right). \]

Thus, from the ratio \( \Delta R_{ex}/R_{ex} \), the exchange rate \( k_{ex} \) can be derived and a correlation plot of \( R_{ex} \) vs. \( \Delta R_{ex} \) shows a linear dependence for a given \( k_{ex} \). Given \( k_{ex} \), the value of \( p_A p_B \Delta_s^2 \) can be derived from \( \Delta R_{ex} \) and/or \( R_{ex} \). Finally, a lower estimate of \( \Delta_{ex} \), \( \Delta_{ex}^{\text{lin}} \), can be obtained, because \( p_A p_B \) has a maximum at \( p_A = 0.5 \).

Ishima and Torchia (34) derived a simple function, which approximates \( R_{ex} \) for fast exchange as well as slow exchange with a skewed population (\( p_A \gg p_B \)):

\[ R_{ex} = \frac{p_A p_B \Delta_s^2 \omega_1^2}{k_{ex}} \left( 1 + \omega_2/k_{ex} \right)^{-1} \]

where, \( \omega_{\text{eff}}(\delta_{1,2}) = \sqrt{3}\beta_{1,2} \). The ratio of \( \Delta R_{ex}/R_{ex} \) is then given by:

\[ \frac{\Delta R_{ex}}{R_{ex}} = \left( \frac{\omega_2}{\omega_1} - \frac{\omega_2}{\omega_2} \right) \left( 1 + \omega_2/k_{ex} \right)^{-1}. \]

Given \( \delta_{1,2} \) and a reasonable maximum of \( \Delta_{ex} \) (\(-6 \) ppm), a correlation plot of \( R_{ex} \) vs. \( \Delta R_{ex} \) calculated using Eq. 6 shows essentially the same linear dependence for \( k_{ex} \) values as when Eqs. 3–5 are used. The \( k_{ex} \) derived from \( R_{ex} \) vs. \( \Delta R_{ex} \) via either set of equations is essentially the same in the fast exchange region. Most importantly, as for Eqs. 3–5, the slope increases with increasing \( k_{ex}^{-1} \) (see below; Fig. 7). Consequently, the \( k_{ex} \) values calculated in this way (Eqs. 6c and/or 5), establish whether fast exchange applies or not without prior knowledge of the exchange timescale. A further note of importance is that for the CPMG settings used, \( R_2(\delta_i) \) is measured in the high \( \omega_{\text{eff}} \) limit (\( \omega_{\text{eff}} > p_A k_{ex} \omega_i \)), so that \( R_{ex} \) is proportional to \( B_0^2 \) independent of whether the exchange is fast or slow to a skewed population (33,34). In conclusion, residues with fast exchange can be identified from the combination of \( R_{ex} \) and \( \Delta R_{ex} \) using either the fast exchange equations or the general fast/slow exchange equations derived by Ishima and Torchia (34).

\( \Delta R_{ex} \) can also be measured at different temperatures (e.g., at T1 and T2), which can be used to derive estimates of the activation enthalpy, \( \Delta H^\theta \), for the exchange process (vide infra). Based on the Boltzmann distribution, the relative populations in an exchanging system do not significantly change upon a small temperature change. Therefore, the difference in \( \Delta R_{ex} \), \( \Delta \Delta R_{ex} \), when the temperature is lowered becomes:

\[ \Delta \Delta R_{ex} = p_A \Delta_s^2 \omega_i^2 \left( \frac{1}{k_{ex,T2}} \left( \frac{\tan(h(k_{ex,T2}\delta_2) - \tan(h(k_{ex,T2}\delta_1)}{k_{ex,T2}\delta_2} - \frac{\tan(h(k_{ex,T2}\delta_1))}{k_{ex,T2}\delta_1} \right) - \frac{1}{k_{ex,T1}} \left( \frac{\tan(h(k_{ex,T1}\delta_1) - \tan(h(k_{ex,T1}\delta_2))}{k_{ex,T1}\delta_1} - \frac{\tan(h(k_{ex,T1}\delta_2)}{k_{ex,T1}\delta_2} \right) \right). \]

From \( \Delta \Delta R_{ex} \), the temperature-induced change in \( k_{ex} \) (\( f = k_{ex,T2}/k_{ex,T1} \)) can be derived, given \( k_{ex} \) and \( p_A p_B \Delta_s^2 \) (established from \( R_{ex} \) and \( \Delta R_{ex} \) at temperature T1). When \( k_{ex}\delta_1 > 3 \) (here when \( k_{ex}\delta_1 < 70 \mu s \), Eq. 7 simplifies, and \( f \) can be calculated from the ratio of \( \Delta \Delta R_{ex} \) and \( \Delta R_{ex} \):

\[ \Delta \Delta R_{ex} \approx -p_A \Delta_s^2 \omega_i^2 \frac{0.875}{k_{ex,T1}} \left( \frac{1}{k_{ex}} - 1 \right)^{-1} = -\Delta R_{ex,T1} \left( \frac{1}{k_{ex,T1}} - 1 \right). \]

According to the transition-state theory in thermodynamics (35,36,37), the rate constant, \( k_{ex} \), is given by:

\[ k_{ex} = \frac{kT}{h} \exp(\Delta G^\theta/RT) = \frac{kT}{h} \exp(\Delta s^\theta/R) \exp(-\Delta H^\theta/RT). \]

Here, \( \Delta G^\theta, \Delta H^\theta, \) and \( \Delta s^\theta \) are the activation free-energy, -enthalpy, and -entropy, respectively; \( k \) is the Boltzmann factor, \( h \) is Planck’s constant, and \( R \) the gas constant. The ratio \( f \) of the exchange rates at two different temperatures then becomes:

\[ f = k_{ex}(T2)/k_{ex}(T1) = \frac{T2}{T1} \exp(-\Delta H^\theta/RT1 T2) \approx \exp(-\Delta H^\theta/RT1 T2). \]

The ratio \( f \) is usually dominated by the exponential factor, so that the right-hand term in Eq. 10 is a good approximation. Hence, the ratio \( f \) can be used to estimate the energy barrier between the exchanging states.

RESULTS

CaM R1 and R2 relaxation rates were measured at 600 and 400 MHz \( ^1H \) frequency and the \( ^15N-\{^1H \} \) NOE data at 600 MHz. The R1, R2 and \( ^15N-\{^1H \} \) NOE data are presented in Fig. 2 and in Supplementary Material (Table S1). In total, 116 residues were analyzed using the PINATA method (22), which is particularly suited for proteins that exhibit extensive nanosecond-timescale internal motions. The data are presented following the flow diagram of the PINATA script.

Identification of nanosecond-timescale internal motion

The presence of nanosecond-timescale motion can be directly identified from a plot of the normalized ratio of \( R_1 \) values measured at two magnetic fields (\( RT_i^\theta \)) versus NOE (22). Fig. 3 shows the theoretical \( RT_i^\theta \) curves for a molecule with a correlation time, \( \tau_i^m \), of 10 ns with one internal motion, \( \tau_i^c \), ranging between 20 ps and 6 ns, and squared order parameter, \( S_i^2 \), ranging between 0.4 and 1.0 (dashed \( \tau_i^c \) contours and solid \( S_i^2 \) contours). The same figure also shows the \( S_i^2 \) contours when an additional nanosecond-timescale internal motion is present with time constant \( \tau_i^s = 2 \) ns and order parameter \( S_i^2 = 0.8 \) (dotted red contours). The interpretation of the \( RT_i^\theta \) versus NOE plots is straightforward.
The $RT_1^f$ value is always 1 in the absence of internal motion or when the internal motion is faster than $\sim 200$ ps. $RT_1^o$ only decreases below 1 when the internal motion is slower than 200 ps or when an additional internal motion with a timescale slower than 200 ps is present. Thus, the presence of nanosecond-timescale internal motion is directly evident from the observation that $RT_1^f$ is smaller than a critical value determined by the experimental error in the $R_1$ measurements. Note that potential variation of the $^{15}N$ CSA between $-150$ and $-200$ ppm hardly affects $RT_1^o$ ($\pm 3\%$), and thus does not affect the conclusions (22).

The $RT_1^f$ values for the CaM:SEF2-1mp complex are superimposed onto the $S^2$ contours in Fig. 3. As can be seen, the $RT_1^f$ values spread around $RT_1^f \approx 0.87$, showing that most residues are affected by the same contribution of nanosecond-timescale motion. This motion may or may not be superimposed onto varying degrees of picosecond-timescale motion. That a two-contribution model is needed follows from a comparison of average experimental and theoretical $R_1$ values. The theoretical $R_1$ values, calculated for a one-contribution model with $S^2$ and $\tau_1$ values consistent with the $RT_1^o$ of 0.87 and varying NOE values (Fig. 3) are always too high, meaning that a two-contribution model needs to be considered. For parameter values of $\tau_{ex} = 2.5$ ns, $S_n^2 = 0.75$, $\tau_{if} = 0.02$ ns, $S_i^2 = 0.80$, and $\tau_m^o = 10$ ns, an $R_1$ value of 1.42 s$^{-1}$ is obtained, which is very close to the average experimental $R_1$. Thus, in addition to the usual picosecond-timescale motions of varying contributions, a small amplitude nanosecond-timescale motion is present for all CaM residues in the CaM:SEF2-1mp complex.

**Determination of the rotational correlation times**

A reliable estimate of the overall rotation correlation times is very important for the further analysis of the relaxation data. In addition, it provides important structural information. The ratio of $R_2$ over $R_1$ ($R_2$ corrected for $R_{ex}$; vide infra) at...
a magnetic field $B$ gives the apparent correlation time, $(\tau_{m}^{s})_{app}$ (Eq. 1). In the absence of internal motion or when there is only a small degree of fast ($<200$ ps) internal motion present, $(\tau_{m}^{s})_{app}$ is equal or close to the true rotation correlation time $(\tau_{m}^{s})$. However, in the presence of nanosecond internal motion, $(\tau_{m}^{s})_{app}$ can be substantially smaller than $(\tau_{m}^{s})$ (22,32). Thus, due to the presence of nanosecond-timescale internal motion, the experimental $(\tau_{m}^{s})_{app}$ for CaM:SEF2-1mp are smaller than the true correlation time, $(\tau_{m}^{s})$. As described elsewhere (22), $(\tau_{m}^{s})_{app}$ can be corrected for the presence of internal motions up to $\sim4$ ns with good accuracy ($\pm0.5$ ns). We note that the potential variation in $^{15}$N CSA does not affect the final corrected $(\tau_{m}^{s})_{app}$ values (22). The $(\tau_{m}^{s})_{app}$ of the CaM:SEF2-1mp complex corrected in this way are shown in Fig. 4a.

Rotational diffusion anisotropy and global orientation

A cylinder-shaped model with a small degree of anisotropy has, according to Eq. 2, a $(\tau_{m}^{s})_{app}$, that depends on the angle, $\Phi$, between the relaxation vector and the long axis of the diffusion tensor. Thus, $(\tau_{m}^{s})_{app}$ contains information about the orientation of the N-H N bond vector via the angle $\Phi$. From the distribution of $(\tau_{m}^{s})_{app}$ values, the shape of the diffusion tensor can be determined (30), whereas the degree of anisotropy can be gauged from the maximum and minimum $(\tau_{m}^{s})_{app}$ values. Given the anisotropy and the maximum $(\tau_{m}^{s})_{app}$, $(\tau_{m}^{s})_{app}$, the angle $\Phi$ that each N-H N bond vector makes with the main axis of the diffusion tensor can be calculated. Moreover, the N-H N bond vectors of an $\alpha$-helix are nearly parallel (within 15°) to the axis of an $\alpha$-helix. Thus, the N-H N bond vectors within an $\alpha$-helix must have similar $(\tau_{m}^{s})_{app}$, and the average $(\tau_{m}^{s})_{app}$ for each helix can be taken to reduce the error. The angle that a helix axis makes with the main axis of the diffusion tensor can therefore be determined with reasonable accuracy from $(\tau_{m}^{s})_{app}$ averaged over a helix.

The variation in $(\tau_{m}^{s})_{app}$ in Fig. 4a shows that the CaM:SEF2-1mp complex tumbles anisotropically. From the estimated maximum and minimum values of $(\tau_{m}^{s})_{app}$, 12.4 ± 0.4 ns and 7.7 ± 0.3 ns, respectively, the anisotropy of the diffusion tensor is calculated to be $2.2 \pm 0.2$ (the average and standard deviation of the 10 highest and 10 lowest $(\tau_{m}^{s})_{app}$). Given an anisotropy of 2.2 ($\Delta = 1.2$) and $(\tau_{m}^{s})$ of 12.4 ns, the angles $\Phi$ that the helices in CaM make with the main axis of the diffusion tensor were derived using Eq. 2: helix I 135 ± 8° (10.5 ± 0.6), helix II 40 ± 3° (10.0 ± 0.2), helix III 32 ± 9° (10.2 ± 0.7), helix IV 44 ± 10° (9.7 ± 0.6), helix V 37 ± 10° (10.3 ± 0.4), helix VI 45 ± 3° (9.6 ± 0.2), helix VII 31 ± 14° (10.6 ± 0.8), and helix VIII 71 ± 14° (8.1 ± 0.5). Here, the average $(\tau_{m}^{s})_{app}$ values for the $\alpha$-helices are given in parentheses and the error on $\Phi$ was based on a uniform error in $(\tau_{m}^{s})_{app}$ of 1.2 ns. A minimum estimate of the anisotropy of 1.6 ± 0.3 is achieved when only the average $(\tau_{m}^{s})_{app}$ in the helices are considered. The $\Phi$ angles, recalculated with the smaller anisotropy, were essentially within the error margins of the earlier estimates. We therefore conclude that not only the pattern of $\Phi$ angle values is reliable, but that also the values are correct within the error margins.

The backbone structures of the CaM N- and C-terminal domains within the CaM:SEF2-1mp complex are expected to be very similar to those in other CaM structures. This is based on the relative small differences in chemical shifts between free and SEF2-1mp bound CaM and the low root mean square deviation of 1.6 Å when the backbones from N- and C-terminal domains from eight other CaM molecules were compared (data not shown). Hydrodynamic calculations on the CaM:SMLCK complex showed that its diffusion tensor is nearly axially symmetric with a $(\tau_{m}^{s})_{app}$ pattern for the $\alpha$-helices that is almost a mirror image of that in the CaM:SEF2-1mp complex, Fig. 4b. This means that the orientation of the diffusion tensor in the CaM:SEF2-1mp complex has rotated by $\sim90°$ relative to that of the CaM:SMLCK complex. In fact, the pattern of $(\tau_{m}^{s})_{app}$ back calculated from the CaM:SMLCK complex when the diffusion tensor would be oriented along an axis perpendicular to the real diffusion tensor closely matches the pattern of $(\tau_{m}^{s})_{app}$ found for the dimeric CaM:SEF2-1mp complex, Fig. 4c.

The qualitative structural implication is, therefore, that the second CaM monomer in the dimeric complex is placed along this axis in such a way that the tips of the N- and C-terminal domains touch each other. This $\sim90°$ rotation of the diffusion tensor together with the relatively small structural changes within each CaM domain puts direct restraints on the overall structure of the CaM:SEF2-1mp complex, which allows us to choose one of the two possible schematic structure models proposed in our earlier study (11) (see Figs. 1 and 4a, right).

Model selection, order parameters, and timescales for internal motion

Based on the qualitative analysis of the RT1 versus NOE plots, Fig. 3, we tested internal motion models with either one (M1; $\tau_{i}$ and $S_{i}$ fitting parameters) or two contributions (M2; $\tau_{i}$, $S_{i}^{2}$, $\tau_{s}$, and $S_{s}^{2}$ fitting parameters) to the internal motion. The $S_{s}^{2}$ was either kept at a uniform constant value (M2I) or optimized together with $\tau_{s}$ and $S_{s}^{2}$ (M2II). A complete overview of the different fit results is given in the Supplementary Material (Tables S3–S8).

As expected from Fig. 3, there is a statistically significant improvement in $\chi^2$ for the vast majority of CaM N-H N backbone vectors when a second internal correlation time is introduced. When different $\tau_{s}$ values were tested, the lowest $\chi^2$ values and hence the best overall fit was found for $\tau_{s}$ equal to 2.5 ns. Although the differences in the $\chi^2$ residuals between M2II models with different $\tau_{s}$ are not statistically significant, it is safe to conclude that there is a slow timescale motion present with a correlation time of roughly 2.5 ns. The
optimized $S^2$ values have an average of 0.75 ± 0.07, and are fairly uniform throughout the sequence, except for five outliers with $S^2$ values between 0.9 and 1. The $S^2$ and $\tau_{\text{off}}$ values extracted from M2II are presented in Fig. 5 and vary around 0.8 and between 5 and 200 ps, respectively. These values are very similar to those found in the core of small well-structured proteins.

In conclusion, the relaxation data show that the dimeric CaM:SEF2-1mp complex has an additional nanosecond-timescale internal motion ($\sim$2.5 ns) superimposed onto the
picosecond-timescale internal motion usually found in well-structured proteins.

**Structural interpretation of the nanosecond-timescale internal motion**

The dimeric CaM:SEF2-1mp complex can in principle show internal modes of motion involving either the individual N- and C-terminal domains and/or the CaM monomers as a whole. Hydrodynamic calculations can provide some indication as to the structural assignment of the internal motions identified from NMR relaxation experiments (see Supplementary Material). Briefly, we find: i), the observed 2.5-ns internal motion in the CaM:SEF2-1mp complex can be attributed to motions of the separate N- and C-terminal domains of CaM, because the free domains have an overall correlation time of ~3 ns. ii), It cannot be excluded, but also not definitely confirmed, that some degree of CaM “monomer” internal motion of ~7–9 ns (estimated tumbling time of free CaM monomer) is also present in the complex.

**Dynamics on micro- to millisecond timescale**

Three different approaches were used to determine which CaM residues in the CaM:SEF2-1mp complex are affected by conformational exchange. Firstly, $R_{ex}$ was derived from the ratio of $R_2$ values measured via a CPMG experiment at two different magnetic fields using the PINATA method. Secondly, the relaxation-compensated CPMG experiment (20) was employed at 500 and 600 MHz to measure $\Delta R_{ex}$. Thirdly, the relaxation-compensated CPMG experiment at 600 MHz was repeated at a lower temperature (300 K instead of 308 K) to measure the change in $\Delta R_{ex}$. $\Delta R_{ex}$ values are from a fit using the two-contribution model, M2II (see text). The time constant $\tau_a$ of the slower internal motion was set to 2.5 ns in the fit. The order parameter of this internal motion, $S^2$, was found to be $0.75 \pm 0.07$ on average. The exact $S^2$, $\Delta S^2$, $\tau_a$, and $\tau_b$ values are given in Supplementary Material (Table S5).

**FIGURE 5** The order parameter, $S^2$ (a) and time constant $\tau_a$ (b) of the fast (picosecond) internal motion. The data are from a fit using the two-contribution model, M2II (see text). The time constant $\tau_a$ of the slower internal motion was set to 2.5 ns in the fit. The order parameter of this internal motion, $S^2$, was found to be $0.75 \pm 0.07$ on average. The exact $S^2$, $\Delta S^2$, $\tau_a$, and $\tau_b$ values are given in Supplementary Material (Table S5).
Table 1) show a weak decrease in exchange rate upon change in temperature \(t \approx 0.9 \pm 0.1\). Consequently, they have a lower activation enthalpy of \(-3.5 \pm 1.3\) kcal/mol. Interestingly, the residues with \(f = 0.4-0.7\) are all located in \(\alpha\)-helices, whereas the residues with \(f \approx 0.9 \pm 0.1\) are all in nonhelical regions except residues 72, 73, and 75, which are part of the less well-defined C-terminal part of helix IV. Thus, the residues with significant exchange broadening can be placed into two groups: those located in helices with a relatively high activation barrier, and those located in loops with a relatively low activation barrier. Akke and co-workers (38) have also found larger \(\Delta H^f\) values for residues that are part of regular secondary structure and smaller values for loop residues, in their study on the C-terminal domain of CaM.

Deriving the activation barriers assuming a two-state exchange is likely to be a significant simplification, as the
conformational landscape may be much more complex. For instance, in the so-called rugged landscape $\Delta H^p$ should be viewed as an average (38). Nevertheless, the trends remain correct, but the $\Delta H^p$ numbers should be considered with proper care.

Nuclei that are influenced by chemical exchange have been shown to correlate with residues known to be critical for protein interactions and enzymatic activity (39,40, 41,42,43,44). We find that residues with significant exchange broadening (Table 1) group in a model structure of the CaM:SEF2-1mp complex into three regions: i), the CaM dimerization interface, i.e., at the tips of the N- and C-terminal domains, ii), close to the hinge region connecting the N- and C-terminal domains, iii), close to the hydrophobic pockets in the N- and C-terminal domains where the target binds (see Fig. 8). This clustering suggests a direct relation between the observed conformational exchange and the CaM:SEF2-1mp interactions (vida infra).

**DISCUSSION**

The dimeric CaM:SEF2-1mp complex shows unique dynamical behavior. Its apparent overall correlation time is $\sim$10 ns at 308 K, as expected for a highly dynamic complex of this shape and size. The complex shows $\sim$20-ps-timescale fast internal motion with order parameters $S_2^2$ around 0.8, as usually seen in well-structured proteins. Interestingly, all N-H vectors are also affected by an additional internal motion of $\sim$2.5 ns with $S_2^2$ of $\sim$0.75. This timescale corresponds well with the expected motions of the N- and C-terminal CaM domains. Thus, the data are consistent with a motional model in which the two N-terminal and two C-terminal CaM domains in the dimeric CaM:SEF2-1mp complex undergo a small-scale wobbling motion with a half-angle of $\sim$20° as estimated from $S_2^2 = 0.75$. However, the present data cannot exclude internal motion involving reorientation of the CaM monomers, which have an excepted timescale of $\sim$7–9 ns. Furthermore, several residues in CaM also undergo conformational exchange on a timescale of $\sim$50 $\mu$s. All the significantly exchanging residues are clustered in well-defined regions, namely on the CaM:SEF2-1mp interaction interface, in the CaM:CaM dimerization interface or in the flexible hinge region that connects the N- and C-terminal domains within the CaM monomer.

The dimeric SEF2-1mp peptide binds within the interior of the eight-shaped CaM dimer via multiple binding alternatives. This is based on the observations that the bound SEF2-1mp lacks any well-defined conformation, and that not all hydrophobic peptide residues simultaneously interact with the hydrophobic patches on the inner surface of the CaM dimer (11). The CaM dimer is created by the tip of the N-terminal domain of one CaM monomer contacting the tip of the C-terminal domain of the other CaM. Hence, there are four hinge regions in the CaM dimer. Two of the hinge regions are formed by the flexible part of the central helix of the CaM molecules, and the other two hinge regions are
formed by the flexible dimerization interfaces. The four hinge regions create a flexible CaM dimer that can easily slide along or wobble over the peptide sequence allowing it to interact with the different binding sites in the interior of the CaM dimer. The sliding process is likely to occur on a nanosecond timescale, although CaM:SEF2-1mp interactions may slow down this process in some instances. Indeed, apart from the nanosecond-timescale wobbling motion of the CaM domains and/or monomers, conformation exchange on the microsecond timescale (∼50 μs) is also evident.

We have compared the characteristics of the CaM:SEF2-1mp interaction with those of other CaM:protein complexes and other protein:protein complexes. In the common wraparound CaM:target binding the hydrophobic patches of CaM and target fit well, resulting in an induced-fit binding where only one binding alternative is needed. Furthermore, the nanosecond-timescale domain motion, present in free CaM, seems to be frozen out, and as far as we know, microsecond-timescale motions are not present in the wraparound binding (10). Note, however, that NMR relaxation data on various protein:target complexes do show a redistribution of motions upon specific target binding. For instance, the barnase (45) and oxalocrotonate tautomerase (46) bind specifically to their rigid target to form a specific complex with essentially one overall conformation. The PLC-γ1 C-terminal SH2 domain (19) and Csk SH3 domain (47) bind specifically to their flexible (peptide) target to form a specific complex, again with essentially one overall conformation. The mouse major urinary protein:pheromone interaction (48) and the topoisomerase I domain interaction with single-stranded DNA (49) have, or suggest to have, the aspect of multiple-binding alternatives in the bound state. However, these binding modes are essentially different from those of CaM:SEF2-1mp, because the target-binding proteins do not undergo large-scale motion in the bound state. Processive enzymes, when sliding along their targets (50), have the closest similarity to CaM:SEF2-1mp binding. However, the sliding mechanism is usually a nonspecific target interaction, whereas in CaM-SEF2-1mp it is specific.

In contrast to known protein:target binding modes, the CaM:SEF2-1mp complex shows large-scale (domain) dynamics in the bound state. The interaction is clearly different from the well-known categories of specific binding by lock-and-key or induced-fit binding, in which the complex in the bound state becomes essentially locked in one conformation. Two aspects of this type of binding as compared to rigid binding are important to discuss, the presence of fast exchange, and the effect on high affinity and specificity.

Fast exchange on the NMR timescale indicates off rates \( k_{\text{off}} > 2 \Delta \delta_N \), which for the CaM:SEF2-1mp complex is \( \sim 30 \text{ s}^{-1} \) (Table 1). Therefore, a purely diffusion-controlled on-rate \( k_{\text{on}} \) of \( 10^8 \text{ s}^{-1} \text{ M}^{-1} \) would put a lower limit on the CaM:SEF2-1mp affinity, \( k_{\text{off}}/k_{\text{on}} > 10^{-6} \). However, in the CaM:SEF2-1mp complex, the two basic peptide arms are trapped in the hydrophobic interior of the CaM dimer. Consequently, the on-rate of the equilibrium between free and CaM-bound peptide will be faster than diffusion controlled, which explains the fast exchange and the high affinity observed in the CaM:SEF2-1mp interaction (11). It is tempting to speculate that the CaM dimer opening-closing process is coupled with the CaM:SEF2-1mp interaction, i.e., when the CaM:SEF2-1mp interaction brakes, one side of the CaM dimer interaction also momentarily brakes. However, due to the close proximity of all involved components, a new CaM:SEF2-1mp interaction is rapidly reformed, which in turn forces the CaM dimer to close again. This collective process could explain the fact that essentially all resonances affected by conformational exchange have approximately the same time constant and are clustered in regions important for either CaM:CaM or CaM:SEF2-1mp interactions.

Mutation studies on E12 (a bHLH protein similar to SEF2-1) have shown a gradual decrease in CaM affinity upon each mutation of an interacting residue, and not the on-off behavior usually seen with lock-and-key or induced-fit binding mechanisms (15). In other words, the CaM:SEF2-1mp binding shows many weak interactions rather than a few strong interactions as found in lock-and-key or induced-fit binding. Thus, removing an interaction is indeed expected to have only a small effect on the affinity. The presence of exchange between multiple-target sites and flexibility in the bound state leads to an important advantage over rigid binding, namely that the affinity (and specificity) is finely tuned. However, this same aspect leads to a lower affinity and specificity than for (cooperative) rigid binding (with all target sites simultaneously and rigidly interacting). This loss is compensated via multimerization. In CaM:bHLH interactions two flexibly bound complexes form a flexibly bound dimer of complexes, defining a geometric context and thereby increasing the number of interactions and thus leading to high affinity and specificity.

In conclusion, the CaM:SEF2-1mp binding seems indeed distinct from other known protein:target binding modes. Its characteristic features are: i) specific high-affinity binding in the presence of fast exchange, ii) accommodation of a relatively nonmatching and flexible target via exchange between multiple binding alternatives, iii) dimerization of CaM upon target binding, and iv), large-scale (domain) motions on the nanosecond and microsecond timescale. Thus, the dynamic features presented here support the conclusion that the conformationally heterogeneous SEF2-1mp, trapped inside the CaM dimer, constantly exchanges between different binding sites. Nature has thus found a way for CaM to specifically recognize a relatively ill-fitting target in its regulation of bHLH transcription factors.

SUPPLEMENTARY MATERIAL

An online supplement to this article can be found by visiting BJ Online at http://www.biophysj.org.
REFERENCES


