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Coupling of Ligand Binding and Dimerization of Helix-Loop-Helix Peptides: Spectroscopic and Sedimentation Analyses of Calbindin D_{9k} EF-Hands

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Isolated Ca2+-binding EF-hand peptides have a tendency to dimerize. This study is an attempt to account for the coupled equilibria of Ca²⁺-binding and peptide association for two EFhands with strikingly different loop sequence and net charge. We have studied each of the two separate EF-hand fragments from calbindin D_{9k} . A series of Ca²⁺-titrations at different peptide concentrations were monitored by CD and fluorescence spectroscopy. All data were fitted simultaneously to both a complete model of all possible equilibrium intermediates and a reduced model not including dimerization in the absence of Ca2+. Analytical ultracentrifugation shows that the peptides may occur as monomers or dimers depending on the solution conditions. Our results show strikingly different behavior for the two EF-hands. The fragment containing the N-terminal EF-hand shows a strong tendency to dimerize in the Ca2+-bound state. The average Ca2+-affinity is 3.5 orders of magnitude lower than for the intact protein. We observe a large apparent cooperativity of Ca2+ binding for the overall process from Ca²⁺-free monomer to fully loaded dimer, showing that a Ca2+-free EF-hand folds upon dimerization to a Ca²⁺-bound EF-hand, thereby presenting a preformed binding site to the second Ca²⁺-ion. The C-terminal EF-hand shows a much smaller tendency to dimerize, which may be related to its larger net negative charge. In spite of the differences in dimerization behavior, the Ca2+ affinities of both EF-hand fragments are similar and in the range $\lg K = 4.6-5.3$. Proteins 2002;47:323-333. \odot 2002 Wiley-Liss, Inc.

Key words: thermodynamic linkage; associationdriven folding; four-helix bundle

INTRODUCTION

The Ca^{2+} ion is an important messenger used in many signalling pathways in biological systems, depending on specific Ca^{2+} -binding proteins. Common to many of these proteins is a Ca^{2+} -binding motif called EF-hand, with a characteristic helix-loop-helix fold. The Ca^{2+} -ion is coordinated by carbonyl and carboxyl oxygens, often complemented with a water molecule. EF-hand motifs normally occur in pairs with the two Ca^{2+} -binding loops coupled through a short antiparallel β sheet. A pair may constitute

a globular domain or be part of a larger entity. Pairing of sites allows for cooperative Ca2+-binding, which enables Ca²⁺-controlled activities to turn on or off over a narrow range of free Ca²⁺ concentration. The EF-hand motif was first discovered in the crystal structure of parvalbumin,1 and over 30 distinct subfamilies of EF-hand proteins have been described.² The subfamilies are often divided into two groups according to the effect of Ca2+-binding on the protein structure. The Ca²⁺-sensor proteins, such as calmodulin and troponin C, are inactive at low Ca²⁺ concentrations $(10^{-7}-10^{-8}\,\mathrm{M})$ and undergo substantial conformational changes when Ca2+ is bound at high Ca2+ concentrations $(10^{-5}-10^{-6} \text{ M})$, leading to activation of target enzymes and a cellular response. The conformational effect of Ca²⁺-binding is much smaller in the other group containing proteins involved in Ca2+ buffer and transport functions.

The presence of two or more Ca²⁺-binding sites in one protein with positive cooperativity makes it difficult to assess the site-specific effects of mutations or changes in solution conditions. To overcome this problem and to study the effects of protein truncation, researchers have studied single, isolated EF-hands, obtained either by protein cleavage or by peptide synthesis. In earlier studies, peptides with different lengths^{3,4} and different amino acid sequences^{5–9} have been compared with regard to secondary structure content and apparent Ca2+ affinity. The latter has been extracted from the data in a simplified way, without regard to peptide association. Studies on EF-hand III from skeletal troponin C have expanded our knowledge greatly, since this was the first case in which an isolated EF-hand was shown to dimerize in the presence of Ca²⁺. ^{10–12} The same behavior was later found for several other EF-hands. 13-15 The coupling of dimerization and Ca²⁺ binding makes it necessary to carefully control the peptide concentration when performing Ca²⁺ titration experiments on single EF-hands. If differences in titration behavior are found with different peptide sequences, this

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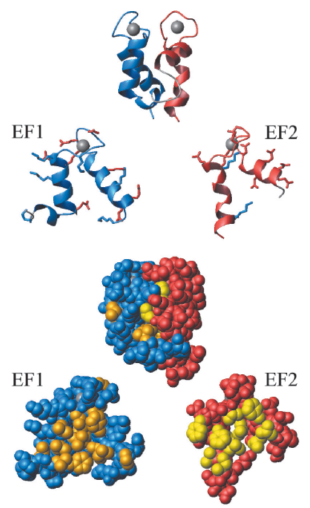


Fig. 1. Crystal structure of Ca^{2+} -saturated calbindin D_{9k} (PDB file 4icb). ⁴³ The intact protein is shown as ribbon and heavy atom representation in the first and third row, respectively, with residues 1–43 in blue and 44–75 in red. The N-terminal EF-hand (EF1) is seen on the left and the C-terminal (EF2) on the right in the second and fourth rows. In rows 2 and 4, EF1 and EF2 are rotated 90° to the left and right, respectively, relative to their orientations in rows 1 and 3, to show the contact surface between the two EF-hands. In the second row are displayed acidic side chains in red and basic side chains in blue. In the third and fourth row are shown hydrophobic side chains in gold (EF1) and yellow (EF2). The picture was generated using MOLMOL. ⁴⁴

may arise from variations in their tendency to self-associate rather than from differences in Ca²⁺ affinity.

We have attempted to resolve the coupled equilibria of $\mathrm{Ca^{2+}}$ binding and dimerization of single EF-hand peptides, using optical spectroscopy to monitor $\mathrm{Ca^{2+}}$ -titration experiments at several peptide concentrations. Sedimentation equilibrium studies were used to assess the oligomerization number of each EF-hand both in the absence and presence of $\mathrm{Ca^{2+}}$, and to study ionic strength effects on peptide association. We have investigated the two EF-hands of calbindin $\mathrm{D_{9k}}$, a small ($\mathrm{M_r}{\sim}8,500$) and very stable for protein in the $\mathrm{Ca^{2+}}$ -buffer/transport group (Figs. 1 and 2). The N-terminal EF-hand (residues 1–43), subsequently called EF1, shows a strong tendency to dimerize when $\mathrm{Ca^{2+}}$ is bound, as well as positive cooperativity of

Fig. 2. Sequence alignment of EF1 and EF2 together with some EF-hands known to form oligomers: s100Bl = N-terminal EF-hand from human S100b protein (apo tetramers). 44 TnC3 = Third EF-hand from chicken skeletal troponin C (Ca²+ dimers). TnC4 = C-terminal EF-hand from rabbit skeletal troponin C (Ca²+ dimers). E6bp4 = C-terminal EF-hand from E6-binding protein (Ca²+ dimers). PV3 = Third EF-hand from silver hake parvalbumin isoform B (Ca²+ dimers). Z is the net EF-hand charge. Ca²+-ligating side chains are shown in bold and the common EF-hand nomenclature on top.

Ca²⁺ binding. The C-terminal EF-hand (residues 44–75), subsequently called EF2, does not dimerize to any great extent under the low-salt conditions, probably due to electrostatic repulsion.

MATERIALS AND METHODS Protein Mutagenesis and Purification

The P43M mutant derived from bovine calbindin D_{9k} was synthesized, expressed in *Escherichia coli*, and purified as described previously. ^{17,18} Agarose gel electrophoresis, SDS-polyacrylamide gel electrophoresis, isoelectric focusing, and ¹H NMR spectroscopy were used to confirm the purity.

The protein is the minor-A form of bovine calbindin D_{9k} with two modifications. The first is a methionine residue added at the N-terminus for expression purposes. The second is the mutation P43M in the flexible loop between the two EF-hands, which makes it possible to cleave it chemically using CNBr.

CNBr Cleavage and Fragment Purification

Cleavage at the methionine residues with CNBr and purification of the resulting fragments with ion-exchange chromatography was performed as described previously¹⁹ with two exceptions: (1) The cleavage was performed on 50 mg of protein and all amounts were scaled accordingly. (2) When eluting the ion-exchange column (DEAE-Sephacel), a linear NaCl gradient from 0.05 M to 0.40 M was used. The purity of the fragments was confirmed by agarose gel electrophoresis.

Agarose Gel Electrophoresis

Agarose gel electrophoresis was carried out in sodium barbitone buffer, pH 8.6, using a 1% agarose gel. Either 2 mM EDTA or 2 mM CaCl₂ was added to the buffer. The protein was visualized by staining with Coomassie blue.

Solutions

All chemicals were of the highest grade commercially available. Lyophilized protein was dissolved in buffer to give protein stock solutions, which were diluted to give the desired protein concentrations. Protein concentrations were confirmed by acid hydrolysis and amino acid analysis. The buffer used was 2 mM Tris/HCl, pH 7.5. ${\rm Ca^{2+}}$ solutions were made from dissolving ${\rm CaCl_2}$ in buffer. EDTA solutions were made by dissolving Titriplex III from Merck (Darmstadt, Germany) in buffer and adjusting the pH to 7.5 with KOH. The residual ${\rm Ca^{2+}}$ contents of the protein fragments and the buffer were determined using a chromophoric ${\rm Ca^{2+}}$ chelator (Quin-2) and UV absorbance spectroscopy.

CD Spectroscopy

The CD spectra were obtained using a Jasco J-720 spectropolarimeter with a Jasco PTC-343 Peltier type thermostatic cell holder. All data were collected at 25.0°C. Quartz cuvettes with path lengths of 1, 2, 4, or 10 mm were used. Only data for which the voltage of the photomultiplier was below 700 V was used (excludes the lower wavelengths at high concentrations due to high absorbance). At higher voltage, the photomultiplier of the instrument becomes saturated and the signal-to-noise ratio of the CD-signal deteriorates rapidly.

Fluorescence Spectroscopy

The fluorescence spectra were obtained using a Perkin Elmer Luminescence Spectrometer LS 50 B connected to a Julabo F25 thermostatic water bath. All data were collected at 25.0°C. The EF1 sample was excited at 274 nm (Tyr) and the EF2 at 260 nm (Phe). The excitation slit was 3 nm and the emission slit 4 nm. Fluorescence cuvettes made of quartz with path lengths of $4{\times}10$ or $10{\times}10$ mm were used.

Analytical Ultracentrifugation

Concentrated stock solutions of the two EF-hands were prepared from freeze-dried material. Samples containing peptide, 2 mM Tris-HCl pH = 7.5, 0.1 M KCl, and either 50μM EDTA or 4 mM CaCl₂ (where specified) were prepared from stock solutions of each individual component. Reference samples were made in an identical fashion excluding the addition of peptide. Samples were analyzed using six channel, 12-mm path length charcoal-filled Epon cells enclosed by quartz windows. Sedimentation equilibrium experiments were performed at 25°C at rotor speeds of 30,000, 40,000, and 50,000 rpm using a Beckman Optima XL-A analytical ultracentrifuge with an An-60Ti rotor. Twenty absorbance scans were averaged. Solution densities were calculated using standard tables listing coefficients for the power series approximation of the change in density $(\Delta \rho)$ for individual components.²⁰ These $\Delta \rho$'s were then summed and added to the density of pure water at 25°C (0.997073 g/mL). Partial specific volumes for both EF-hand fragments were calculated from the weighted average of the partial specific volume of the individual amino acids.²⁰ The Nonlinear Least Squares Program (HID) obtained from the Analytical Ultracentrifugation Facility at the University of Connecticut was used for the curve fitting data analysis.

Analysis of Ca²⁺ Binding Data

The data analysis was carried out using the software MATLAB (The Math Works, Inc.). A MATLAB program was written using the Levenberg-Marquardt algorithm for least square fits.²¹ The program, called LMFIT, allows simultaneous fitting of a large number functions and data sets so that one or more common constants may be determined with higher precision.

To correct for the use of different cuvettes, the CD signals were multiplied by a factor 10/d, where d is the path length in mm. The fluorescence data obtained in a 10×10 mm cuvette was corrected by division of a factor 1.30 to be comparable to the 4×10 mm cuvette data. This factor was obtained from several fluorescence measurement of identical samples in 4×10 mm and 10×10 mm cuvettes. The total peptide concentration, $A_{\rm tot}$, at each titration point was calculated from A_0 (the initial value of $A_{\rm tot}$) and the dilutions due to Ca^{2+} additions. The total Ca^{2+} concentration, $Ca_{\rm tot}$, was calculated from the residual Ca^{2+} contents of the protein fragments and the buffer, the Ca^{2+} additions and the dilutions.

In a preliminary analysis, each titration curve was fitted independently, assuming no dimerization, to a single Ca^{2^+} -binding constant, K_1 . This was done to assess the quality of the data and the effect of protein concentration. This simple model will subsequently be referred to as the 2-state model.

$$A + Ca \stackrel{K_1}{\rightleftharpoons} ACa$$

$$K_1 = \frac{[ACa]}{[A][Ca]} \tag{1}$$

The total ${\rm Ca^{2^+}}$ concentration, ${\rm Ca_{tot}}$, is the sum of the concentrations of all calcium-containing species. In this case

$$Ca_{\text{tot}} = [Ca] + [ACa] \tag{2}$$

The total peptide concentration, $A_{\rm tot}$, is the sum of the concentrations of all peptide-containing species. In this case

$$A_{\text{tot}} = [A] + [ACa] \tag{3}$$

Substitution for [ACa] in eq. (2) using eq. (1) and rearrangement gives

$$[Ca] = \frac{Ca_{\text{tot}}}{1 + K_1[A]} \tag{4}$$

Substitution for [ACa] in eq. (3) using eq. (1) and eq. (4) gives

$$A_{\text{tot}} = [A] + \frac{K_1[A]Ca_{\text{tot}}}{1 + K_1[A]}$$
 (5)

Multiplication with $(1+K_1[A])$ and rearrangement gives

$$K_1[A]^2 + (1 + Ca_{tot}K_1 - A_{tot}K_1)[A] - A_{tot} = 0$$
 (6)

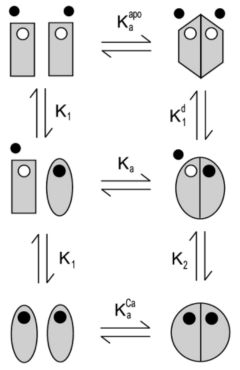


Fig. 3. The 5-state model including all possible dimerization and Ca $^{2+}$ -binding events that can take place with an isolated EF-hand fragment. The 4-state model is obtained by omitting apo dimers and the equilibria described by association constants K_a^{apo} and K_1^{d} .

This quadratic equation may be solved analytically using the standard solution of quadratic equations. Only the positive root is valid, since [A] cannot be negative. The spectroscopic data are fitted using molar spectroscopic values for the apo form and the Ca^{2+} -loaded form: Y_{apo} and $Y_{\mathrm{Ca}}.$

$$Y = Y_{Ca} + \frac{[A]}{A_{\text{tot}}} (Y_{\text{apo}} - Y_{Ca})$$
 (7)

CD and fluorescence spectroscopy are methods where dilution affects the signal and the peptide concentration change were accounted for by multiplying the right hand side of equation (7) with the ratio A_{tot}/A_0 .

The fact that the errors vary in complicated ways with factors such as wavelength, path length, method, and concentration made careful assessment of the measurement errors necessary. The measurement errors were assumed to be absolute in the case of CD spectroscopy and to be relative in the case of fluorescence spectroscopy. To estimate the magnitude of the errors, a scaling factor was allowed to change until χ^2 was equal to the number of degrees of freedom. This was done for each titration separately, using the simple two-state model, and the errors were used in the fittings of the more complicated models.

In a more extensive analysis, a complete model of all possible equilibria as shown in Figure 3 was used. This model will subsequently be referred to as the 5-state model, because the peptide can exist in five different

states: apo monomer, Ca₁-monomer, apo dimer, Ca₁-dimer, or Ca₂-dimer. The corresponding equations are:

$$K_1 = \frac{[ACa]}{\lceil A \rceil \lceil Ca \rceil} \tag{8}$$

$$K_a = \frac{[A_2Ca]}{[A][ACa]} \tag{9}$$

$$K_{2} = \frac{[A_{2}Ca_{2}]}{[A_{2}Ca][Ca]} \tag{10}$$

$$K_a^{Ca} = \frac{[A_2 C a_2]}{[A C a]^2} \tag{11}$$

$$K_a^{\text{apo}} = \frac{[A_2]}{[A]^2} \tag{12}$$

$$K_1^d = \frac{[A_2 Ca]}{\lceil A_2 \rceil \lceil Ca \rceil} \tag{13}$$

Hence, K_a , K_a^{apo} , and K_a^{Ca} are dimerization constants, and K_1 , K_2 , and K_1^d are Ca^{2+} association constants. Since the six equilibria (the two equilibria depicted in the left column of Figure 3 are actually the same equilibrium) are connected in a thermodynamic double cycle, it is only necessary to determine four of them [e.g., eqs. (8), (9), (10), and (12)]. To account for all the Ca^{2+} -bound species present, eq. (2) is modified into eq. (14).

$$Ca_{\text{tot}} = [Ca] + [ACa] + [A_2Ca] + 2[A_2Ca_2]$$
 (14)

Substituting for the species [ACa], [A₂Ca] and [A₂Ca₂] using eqs. (8), (9), and (10) and rearranging we arrive at the following quadratic equation

 $2K_1K_aK_2[A]^2[Ca]^2$

+
$$(1 + K_1[A] + K_1K_a[A]^2)[Ca] - Ca_{tot} = 0$$
 (15)

Eq. (3) is modified to

$$A_{\text{tot}} = [A] + [ACa] + 2[A_2Ca] + 2[A_2Ca_2] + 2[A_2]$$
 (16)

Substitution using eqs. (9), (10), (11), and (13) gives

$$A_{\text{tot}} = K_1[A][Ca] + 2K_1K_a[A]^2[Ca] + 2K_1K_aK_2[A]^2[Ca]^2 + 2K_a^{\text{apo}}[A]^2 \quad (17)$$

[Ca] may be solved analytically from eq. (15). When this expression is used instead of [Ca] in eq. (17), it becomes impossible to solve analytically. Therefore, this is done numerically with a MATLAB command called fzero. This algorithm was originally written by T. Dekker and uses a combination of bisection, secant, and inverse quadratic interpolation methods.²² The various peptide species are assumed to contribute linearly to the CD or fluorescence signal.

$$Y_A = m_A[A] \tag{18}$$

$$Y_{ACa} = m_{ACa}[ACa] \tag{19}$$

$$Y_{A_2Ca} = m_{A_2Ca}[A_2Ca] \tag{20}$$

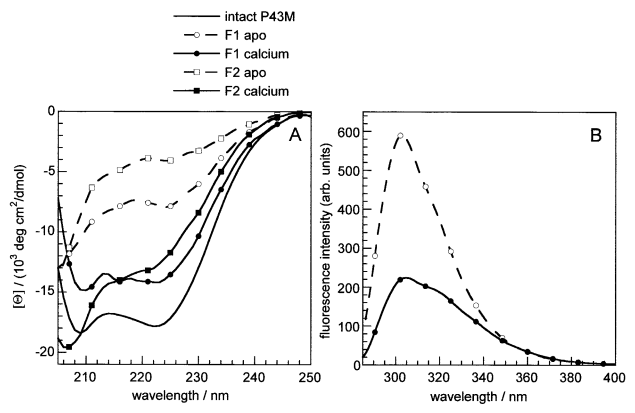


Fig. 4. CD and fluorescence spectra of the isolated EF-hands from calbindin D_{gk} . **A:** The mean residue ellipticity, [Q], for EF1 (30 μ M) and EF2 (43 μ M) as compared to intact calbindin D_{gk} . The same samples were used for both the apo and Ca^{2+} spectra. First, 50 μ M EDTA was added and the apo spectra were obtained. Then 10 mM $CaCl_2$ was added and the Ca^{2+} spectra were obtained. **B:** Fluorescence emission spectra of EF1 (43 μ M) in the absence and presence of 2.5 mM $CaCl_2$.

$$Y_{A_2Ca_2} = m_{A_2Ca_2}[A_2Ca_2] \tag{21}$$

$$Y_{A_2} = m_{A_2}[A_2] (22)$$

The spectroscopic signal is obtained by summation of the contributions of the different peptide species and addition of the background signal, b.

$$Y = Y_A + Y_{ACa} + Y_{A \circ Ca} + Y_{A \circ Ca} + Y_{A \circ} + b \tag{23}$$

Since evidence for the existence of an apo multimeric state has been found in one case only, ²³ attempts were also made to fit the data to a simplified model not including the apo dimer. This model will subsequently be referred to as the 4-state model. Eqs. (16), (17), and (23) were simplified accordingly into eqs. (24), (25), and (26), respectively.

$$A_{\text{tot}} = [A] + [ACa] + [A_2Ca] + [A_2Ca_2]$$
 (24)

$$A = K_1[A][Ca] + 2K_1K_a[A]^2[Ca] + 2K_1K_aK_2[A]^2[Ca]^2$$
 (25)

$$Y = Y_A + Y_{ACa} + Y_{A > Ca} + Y_{A > Ca} + b$$
 (26)

Several data sets were used simultaneously to obtain better precision in the equilibrium binding constants using the multiple function option in LMFIT. For EF1, CD data at different wavelengths were combined with each other and with fluorescence data at different wavelengths by allowing the constants m_A , m_{ACa} , m_{A2Ca} , m_{A2Ca2} , and b to

be wavelength-specific, but keeping K_1 , K_a , and K_2 constant for all wavelengths. For EF2, the data for different concentrations and different wavelengths was combined by assigning individual values of $Y_{\rm apo}$ and $Y_{\rm Ca}$ to each combination of concentration and wavelength, but letting K_1 be common for all data sets. In the text, the binding constants are discussed as $\lg K (\lg K = {}^{10} \log K = -pK)$.

RESULTS

CNBr Cleavage and Isolation of Fragments

The calbindin D_{9k} Pro43 \rightarrow Met mutant (P43M) was cleaved by CNBr to produce the single EF-hand fragments EF1 (residues 1–43) and EF2 (residues 44–75). The two fragments have different net charges (–1 for EF1 and –6 for EF2) and could, therefore, be separated from each other using ion exchange chromatography in the presence of EDTA, followed by desalting on a G25 gel filtration column. The yields of pure desalted fragments were 16–19 mg of EF1 and 8–12 mg of EF2 when 50 mg intact protein was cleaved.

CD Spectra

The CD spectra of the two EF-hand fragments of calbindin D_{9k} in the presence and absence of Ca^{2+} are seen in Figure 4(A) together with the CD spectrum of the intact protein. The CD signals of both EF1 and EF2 change

		Rotor speed (rpm)					
Sample conditions	30,000	40,000	50,000	Global			
42.6 μM EF2, Apo	$5,000 \pm 600$	$3,\!800 \pm 400$	$2,900 \pm 200$	$3,\!100 \pm 200$			
42.6 μM EF2, Apo, KCl	$11,400 \pm 1,100$	$7,600 \pm 700$	$5,800 \pm 300$	$6,400 \pm 500$			
$42.6 \mu\mathrm{M}$ EF2, CaCl_2	$10,000 \pm 900$	$9,900 \pm 400$	$9,300 \pm 300$	$9,500 \pm 300$			
42.6 μM EF2, CaCl ₂ , KCl	$10,700 \pm 700$	$10,000 \pm 400$	$9,200 \pm 300$	$9,\!500 \pm 300$			

[†]A single species analysis treats the molecular weight as a variable in the curve fitting analysis. The global analysis analyzes the data from the three rotor speeds simultaneously. Molecular mass in Daltons; the theoretical monomer molecular weight is 3,674.0 Daltons.

substantially upon Ca2+ addition, making it possible to use CD spectroscopy as a probe of Ca²⁺ binding. The Ca^{2+} -form of EF1 yields a spectrum typical of an α -helical structure. It closely resembles the spectrum of the intact protein, although the magnitude of the signal is smaller. This is an indication that the Ca2+-form of EF1 has a secondary structure similar to the highly helical calbindin D_{9k}. The reduction in signal compared to the intact protein may be explained by mere fraying of helices, without reduction of the number of helical residues, according to a theoretical study by Hirst and Brooks.²⁴ The CD spectrum of EF1 may be consistent with a symmetric dimer structure like the one found for EF-hand III of chicken troponin C¹². The Ca²⁺-form of EF2 also yields a spectrum with substantial signal intensity, but the ratio of the intensities at 222 and 208 nm is smaller than for EF1, which may suggest a lower degree of α -helical structure. The CD signals of the Ca²⁺-free (apo) forms, especially for EF2, are much weaker than the signals for the Ca²⁺ forms, suggesting that Ca²⁺ stabilizes the secondary structure elements and induces folding.

Fluorescence Spectra

EF1 contains one tyrosine and two phenylalanine residues, and the tyrosine fluorescence spectra in the presence and absence of $\mathrm{Ca^{2+}}$ are shown in Figure 4(B). The spectra reveal a large decrease in fluorescence signal upon calcium binding, which may be monitored during a $\mathrm{Ca^{2+}}$ -titration experiment. Since folding of proteins generally results in decreased amounts of tyrosine emission, 25 the large decrease in fluorescence signal upon $\mathrm{Ca^{2+}}$ binding may be taken as an additional indication of a $\mathrm{Ca^{2+}}$ -induced folding process. EF2 contains only 3 aromatic residues, all of which are phenylalanines. The fluorescence of EF2 in the presence and absence of calcium was investigated, but the signal was overall very weak and the differences between the spectra of the $\mathrm{Ca^{2+}}$ and apo forms were very small (data not shown).

Analytical Ultracentrifugation of EF2

Sedimentation equilibrium analytical ultracentrifugation was performed to probe the homo-oligomerization state of EF2 in both the absence and presence of $\mathrm{Ca^{2^+}}.$ In the absence of calcium, the monomer molecular weight (3,674.0 Da) is underpredicted (Table I).This is likely due to electrostatic repulsion of the highly charged apo form of EF2 (Z = -6). Thus, at 42.6 μM , apo EF2 is completely monomeric with no evidence for self-association. In the

presence of 0.1M KCl, apo EF2 shows significant self-association as reflected in the elevated molecular weight $(6,400\pm500;$ obtained from global fits using all three concentrations simultaneously). The observed trend of decreasing apparent molecular weight with increasing rotor speed, which is an evidence for multiple species, further supports this conclusion.

The global molecular weight of EF2 in the presence of Ca^{2^+} is intermediate between a dimer and trimer. Thus, there is evidence for association above a dimer in the presence of a significant excess of Ca^{2^+} .

Ca²⁺ Titration of EF2

One series of Ca²⁺-titrations was performed on EF2 and monitored by CD at 210, 215, 220, 225, and 230 nm. The 220-nm data is shown in Figure 5(A). EF2 was titrated with Ca²⁺ under a wide range of peptide concentrations $(1.7-250.4 \mu M)$, and attempts to fit the individual titration curves using a single Ca2+-binding constant yielded essentially the same result for all curves. Attempts to fit the titration data to the more complex 4-state and/or 5-state models (Fig. 3) gave no significant improvement over the 2-state model, and all binding constants except K₁ were completely undefined with the errors larger than the obtained values. Therefore, all titrations, including all combinations of concentrations and wavelengths, were fitted simultaneously to a common K1-value using the 2-state model while $Y_{\rm Ca}$ and $Y_{\rm apo}$ were allowed to be titration-specific. Data with total ${\rm Ca^{2+}}$ concentration above 1 mM were discarded before the final fitting because the increased ion strength may affect the equilibria of the highly charged EF2 (Z = -6 in the apo form). The resulting fit is shown as the solid line in Figure 5(A). It yielded the association constant $lgK_1 = 4.620 \pm 0.004$. A deviation from the fitted curves can been seen at the highest EF2 concentrations (166.9 and 250.4 μM), probably due to influence of dimerization at these peptide concentrations. When plotting the degree of Ca²⁺ saturation vs. the free Ca²⁺ concentration [Fig. 5(B)], it is clear that overall the EF2 data is consistent with a single binding constant as the midpoint free Ca²⁺ concentration is the same for all titrations. The value of K₁ corresponds to a dissociation constant of 24.0 \pm 0.2 μ M, which is equal to the midpoint in the transformed binding curves shown in Figure 4(B).

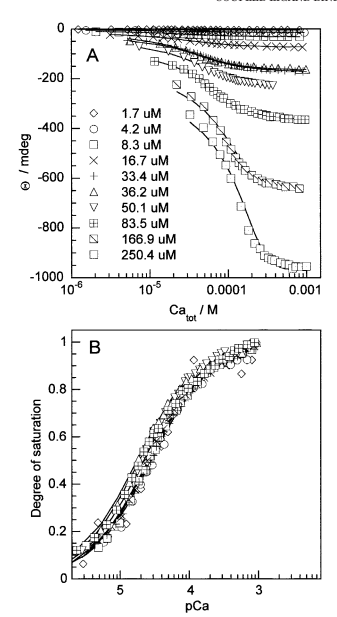


Fig. 5. **A:** Ca^{2+} titrations of EF2 as measured by CD spectroscopy at 220 nm. The solid line was obtained by least-squares fitting simultaneously to all data using the 2-state model. **B:** The data and fitted curve in A for 1.7–83.5 μ M EF2 transformed to the degree of Ca^{2+} saturation as a function of free Ca^{2+} .

Analytical Ultracentrifugation of EF1

Sedimentation equilibrium analytical ultracentrifugation was performed to probe the homo-oligomerization state of EF1 in both the absence and presence of $\mathrm{Ca^{2^+}}$. The apparent molecular weight in the absence of $\mathrm{Ca^{2^+}}$ is higher than that expected for a monomer across the eightfold concentration range studied, from 39 to 312 $\mu\mathrm{M}$ (Table II).However, the apparent molecular weight is independent of concentration, contrary to the results expected for a self-associating system in solution. Additionally, no trend of decreasing molecular weight with increasing rotor speed is observed, suggesting that a single

species is present in solution. Taken together, these results suggest that EF1 is monomeric in the absence of Ca²⁺.

In the presence of 4 mM Ca²⁺, the molecular weight is only slightly elevated above that of a dimer (9,757.2 Da). Modeling higher order association states above that of a dimer did not improve the fit of the model to the data (data not shown). The elevated observed molecular weights in both conditions could be due to ill-determined values for the calculated peptide partial specific volumes or solvent densities. Equally plausible is the possibility that a small amount of irreversible aggregation formed in the concentrated stock solution. Nevertheless, the results strongly correlate with an oligomerization state of a monomer and dimer in the absence and presence of Ca²⁺, respectively.

Ca²⁺ Titration of EF1

Two series of Ca²⁺-titrations were performed on EF1 with peptide concentrations ranging from 1.6 to 86 μM. The first titration series was monitored with CD at 222 nm and is shown in Figure 6(A). The second was monitored with CD at 205, 210, 215, 220, 225, and 230 nm and with fluorescence at 295, 300, 303, 305, 310, 315, 320, 325, and 330 nm. In Figure 6(B), 303-nm fluorescence data are shown as an example. Attempts to fit each individual titration curve to a single apparent Ca²⁺-binding constant did not yield consistent results and instead showed a strong correlation between fragment concentration and apparent Ca²⁺ affinity. Therefore, all titrations were fitted simultaneously allowing for dimerization of the peptides. A fit to the Ca²⁺-titration data using the 4-state (no apo dimers, Fig. 3) yields lgK $_1$ = 4.41 \pm 0.01, lgK $_a$ = 4.67 \pm 0.03, and $\lg K_2 = 4.95 \pm 0.01$. This fit is shown as a solid line in Figure 6(A,B) for the 222-nm CD data and the 303-nm fluorescence data, respectively. The association constant for dimerization f two Ca2+-bound EF1 can be calculated from the three fitted constants as lgK_a(Ca) = $lgK_a + lgK_2 - lgK_1 = 5.21$. The corresponding fit of the 5-state model (apo dimers allowed) looks almost exactly the same to the eye (curve not shown), but a slightly lower χ^2 is found. The results of the analytical ultracentrifugation suggest that Ca²⁺-bound EF1 is dimeric whereas the apo form is monomeric, and therefore supports the 4-state model. For comparison of the titrations at different EF1 concentrations, the degree of saturation was plotted vs. the free Ca²⁺ concentration. The midpoint of the Ca²⁺titration curves is clearly dependent on peptide concentration, as expected from the preliminary fits of the individual titration curves, which yielded a concentration-dependent apparent Ca²⁺ affinity.

Since two independent titration series were available and since for the last titration series, two different methods were used (CD and fluorescence), fitting to different sub-sets of the data gave us an idea of how well determined the different association constants are. Overall, the Ca²⁺-binding constants were better determined than the dimerization constant(s) for both models.

			Rotor speed (rpm)				
Sample conditions	$Peptide\left(\mu M\right)$	30,000	40,000	50,000	Global		
156 μM EF1, Apo	39	$6,\!200 \pm 800$	$6,\!500 \pm 700$	$6,\!200 \pm 400$	$6,200 \pm 400$		
	156	$6,\!500 \pm 600$	$6,600 \pm 500$	$6,400 \pm 200$	$6,400 \pm 300$		
	312	$6,\!300 \pm 800$	$6,900 \pm 700$	$5,500 \pm 600$	$6,100 \pm 600$		
$156 \mu \mathrm{M} \mathrm{EF1}, \mathrm{CaCl}_2$	39	$11,\!200 \pm 100$	$10,\!400 \pm 400$	$10,100 \pm 300$	$10,\!200 \pm 300$		
	156	$11,\!300 \pm 700$	$10,700 \pm 400$	$10,100 \pm 200$	$10,\!300 \pm 200$		
	312	$10,\!000 \pm 100$	$10{,}100\pm700$	$9,\!200 \pm 600$	$9,700 \pm 500$		

[†]A single species analysis treats the molecular weight as a variable in the curve fitting analysis. The Global analysis analyzes the data from the three rotor speeds simultaneously. Molecular mass in Daltons; the theoretical monomer molecular weight is 4878.6 Daltons. All data sets from the three peptide concentrations were analyzed simultaneously. The global molecular weight is not dependent on peptide concentration, thus this analysis is valid.

DISCUSSION

The ability of EF-hand peptides to form homodimers or higher oligomers makes it necessary to analyze ${\rm Ca^{2^+}}$ -binding data in terms of the coupled equilibria of ${\rm Ca^{2^+}}$ -binding and association. Furthermore, the analysis is greatly facilitated if another method, like analytical ultracentrifugation, can provide independent information on the oligomerization number and whether oligomers can be formed both in the absence and presence of ${\rm Ca^{2^+}}$. In the present work, we have attempted to resolve these questions by combining data from analytical ultracentrifugation and ${\rm Ca^{2^+}}$ titrations followed by optical spectroscopy.

EF1

The N-terminal EF-hand in calbindin D_{9k} , EF1, does not agree with the consensus sequence. The loop accommodates two extra residues and Ca^{2+} coordination is mainly via backbone carbonyls. In addition, the net charge of EF1 is low (-1 without Ca^{2+} and +1 with Ca^{2+} bound), although there are several negatively charged residues in and around the Ca^{2+} -binding site. In the intact protein, the N-terminal site is less flexible than the C-terminal site. This is observed as lower rates of backbone fluctuation²⁶ and a stringent selection for Ca^{2+} over other mono-, diand trivalent metal ions^{27,28}.

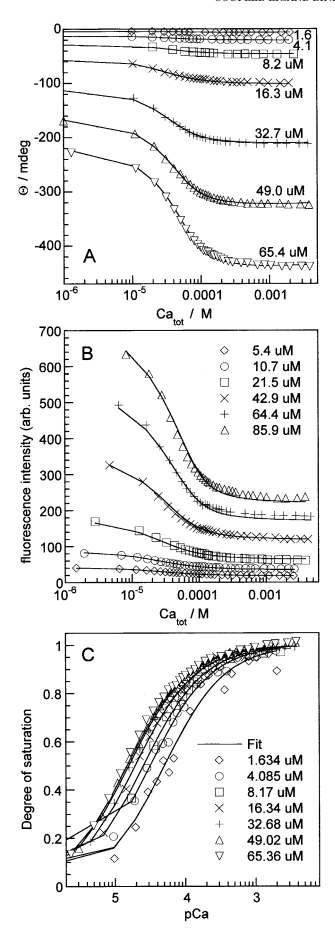
The analytical ultracentrifugation results imply that EF1 is monomeric in the absence of calcium and dimeric in the presence of Ca²⁺. This suggests that the 4-state model, which neglects dimerization in the apo state, is of sufficient complexity for analysis of the Ca²⁺ titrations. We may then conclude that for EF1, the first Ca²⁺ ion binds to a monomer with an affinity of $\lg K_1 = 4.41$. The Ca^{2+} bound EF1 may then associate with another Ca²⁺-bound EF1 with an association constant of $\lg K_a(Ca) = 5.21$. Alternatively, it may associate with an apo EF1 (lg K_a = 4.67), and a second Ca²⁺ ion may bind to the dimer with an affinity of $\lg K_2 = 4.95$. Using the four association constants, the concentrations of the four species EF1 apo, EF1Ca, EF12Ca, and EF12Ca2 may be calculated as a function of total Ca²⁺, as shown in Figure 7 for two different EF1 concentrations (1 and 100 µM). At low Ca²⁺ concentration ($< 1 \mu M$), there is no difference between the two cases, but as Ca2+ is added, the main species formed at 1 μ M EF1 is the Ca²⁺-bound monomer, with little accumulation of EF1₂Ca₂. At 100 μ M EF1, there is an accumulation of an EF1₂Ca intermediate between 10 and 100 μ M Ca²⁺, which is then converted to mainly EF1₂Ca₂ and some EF1Ca.

For EF1, which dimerizes only with calcium bound, it is evident that there is a strong coupling between Ca²⁺ binding, folding, and association of the EF-hand.

EF2

The C-terminal EF-hand in calbindin D_{9k} , EF2, is a standard EF-hand with all the features expected from the consensus sequence. The net charge of EF2 is -6 in the apo state and -4 with Ca^{2+} bound. The C-terminal site uses mainly side-chain carboxylates to coordinate Ca^{2+} , and can therefore adjust the coordination sphere to accommodate a number of different divalent metal ions like Cd^{2+} , Mg^{2+} , Mn^{2+} , and also the trivalent lanthanide ions like Lu^{3+} , Eu^{3+} , and many others.

The apo form of EF2 is clearly a monomer in the absence of added salt. The analytical ultracentrifugation data underpredicts the monomer molecular weight, an indication of measurable electrostatic repulsion. Addition of Ca²⁺ and/or KCl seems to promote dimerization of the peptide. In contrast to EF1, very similar apparent Ca²⁺binding constants were obtained with different concentrations of the EF2 peptide. Hence, under the conditions of the Ca²⁺ titrations, dimerization does not influence the Ca²⁺ affinity of EF2. This may seem unexpected, because the high net negative charge of an EF2 homodimer would provide a very strong electrostatic attraction for Ca²⁺ ions. However, electrostatic repulsion between the two EF2 chains in the homodimer may prevent an optimal arrangement of the two Ca²⁺ loops. In earlier experiments, it was observed that the Ca²⁺ affinity is reduced in a predictable way when the net charge of intact calbindin D_{9k} is changed from -7 to -6, -5, or -4 through site-directed mutagenesis. 29,30 However, when the net charge is changed to -8, as in either of the three single mutants N21D, Q22E, and N56D, the effect is either a decrease, or only a small increase, in Ca²⁺ affinity, ^{31,32} suggesting that when the negative charge becomes too large, structural destabilization leads to effects opposing the affinity gain resulting from an increased electrostatic attraction of Ca²⁺.



Unfavorable electrostatic interactions between EF2 monomers in the apo form seem to prevent dimerization at low ionic strength. As discussed previously, the presence of salt increases the dimerization constant of apo EF2. In the presence of ${\rm Ca^{2+}}$, dimers are observed both in the absence or presence of added KCl. However, the CD spectrum [Fig. 4(A)] suggests that the dimers do not have as well-defined helix-loop-helix structure as the intact protein and ${\rm Ca^{2+}}$ -bound EF1 homodimer.

Cooperativity of Calcium Binding

In a system containing two Ca²⁺-binding sites, there is a possibility for cooperativity between the sites, i.e., binding of the first Ca²⁺-ion may change the affinity for the second ion. Cooperativity is most readily quantitated in a symmetric system with two identical sites. There seems to be no cooperativity in the EF2 system as the data are well fitted by a single apparent Ca2+-binding constant. However, EF1 deserves a closer analysis. If we focus on the consecutive process (Ca2+ binding to monomer)-(dimerization)-(Ca²⁺ binding to the empty second site in the dimer), an apparent cooperativity may be calculated from the ratio of $\rm K_2$ and $\rm K_1$, as $\Delta \Delta G = -RT \ln{(\rm K_2/\rm K_1)} = -3.1$ kJ/mol. There is hence a positive cooperativity ($\Delta\Delta G < 0$), and the second Ca²⁺ ion is bound with a higher affinity than the first $(K_2/K_1 = 3.5)$. The degree of cooperativity is about half of that in the intact protein ($\Delta\Delta G = -8.2 \text{ kJ/mol}^{33}$). The cooperativity observed for EF1 may arise from the fact that Ca²⁺ binding induces folding of the EF-hand (observed by CD spectroscopy), and this is at least partly propagated to its dimerization partner. In general, a large entropic contribution to the Ca²⁺ affinity for a protein or peptide comes from the release of water molecules from the first hydration shell of the solvated Ca²⁺ ion. Ca²⁺ binding is also promoted by electrostatic interactions and preformation of the binding site, while the affinity is reduced by conformational changes driven by the Ca²⁺ binding event (for review see Linse and Forsén³⁴). Binding of the first Ca²⁺ ion to one EF1 peptide promotes dimerization with and folding of another EF1 copy, such that the second Ca²⁺ ion can more or less simply "pop in." While both ions may get all the benefits of water release and interactions in the Ca²⁺ bound state, the first one has to pay most of the penalty of driving the conformational change. The first step will, therefore, display a lower Ca²⁺ affinity than the second, and the result is positive cooperativity.

Comparison to Other EF-Hand Peptides

Ca²⁺ binding data have been reported for a number of single EF-hand peptides, and the apparent affinities are

Fig. 6. Ca^{2+} titrations of EF1. **A:** The first series of titrations as monitored by CD spectroscopy at 222 nm. The solid line was obtained by least-squares fitting simultaneously to all data using the 4-state model. **B:** The second series of titrations as monitored by fluorescence spectroscopy at 303 nm. The solid line was obtained by least-squares fitting simultaneously to all data using the 4-state model. **C:** The data and fitted curve shown in A transformed to degree of Ca^{2+} saturation vs. the free Ca^{2+} concentration, [Ca].

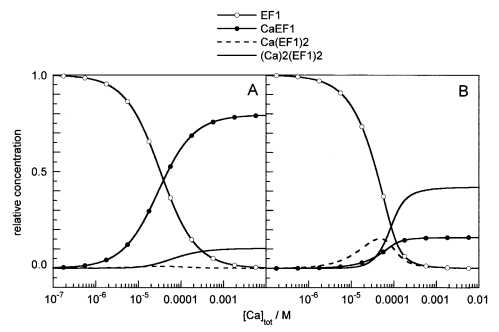


Fig. 7. Relative concentrations of EF1 apo, EF1Ca, EF1 $_2$ Ca, and EF1 $_2$ Ca $_2$ as a function of total Ca $^{2+}$ concentrations, Ca $_{tot}$, calculated using the fitted association constants of the 4-state model for a total EF1-concentration of 1 μ M (**A**) and 100 μ M (**B**).

usually in the 10³–10⁵ M⁻¹ range. Among the more careful studies are those of EF-hand III from troponin C (net charge of -6 in the Ca²⁺ form) in which attempts were made to account for the coupled equilibria of Ca²⁺ binding and dimerization.³⁵ One-dimensional NMR-spectra were obtained during one Ca²⁺ titration experiment with constant peptide concentration and one series of dilution experiments with constant Ca²⁺:peptide concentration ratio; in total 27 data points. Theoretical results for a large number of different combinations of binding constants were calculated and the combination of constant values that yielded the lowest standard deviation between calculated and experimental data was chosen. This procedure yielded association constants of $\lg K_1 = 5.5$ for the first Ca^{2+} and $lgK_a = 5.0$ for the dimerization, but the binding of the second Ca²⁺ ion to the dimer was found to be too weak to be determined. 35 The fact that the affinity for the first Ca²⁺ ion was much higher than for the second implies a strong negative cooperativity in the troponin C EF-hand III system. Clearly, this system behaves differently than both EF1 and EF2 from calbindin D_{9k}.

Comparison to the EF1-EF2 Heterodimer

Although the single EF-hands form homodimers in the presence of ${\rm Ca^{2^+}}$, the affinity between two EF1 peptides, ${\rm lg~K}=5.21$, is more than six orders of magnitude lower than the association constant for formation of heterodimer between EF1 and EF2, ${\rm lg~K}=11.5$. The heterodimerization constant was recently measured using surface plasmon resonance technology. This enormous difference implies a very high specificity for heterodimers over homodimers in an EF1-EF2 mixture. In fact, the two EF-hands interact so tightly that they

resist separation by ion exchange in the presence of Ca^{2+} . ^{19,37} The remarkable specificity for heterodimers is in line with other studies, ^{37–40} and was recently found to allow for dimerization of calbindin D_{9k} via EF-hand swapping. ⁴¹

Comparison to Intact Calbindin

EF-hand sites in intact proteins have affinities spanning a much wider range all the way up to 10⁹ M⁻¹. The Ca²⁺-binding constants obtained for EF1 and EF2 (lg K = 4.4-5.0) are three to four orders of magnitude lower than those measured for intact calbindin D_{9k} (lg K = $7.8-8.6^{29}$) under low ionic strength conditions. Hence, the specific contacts between the two EF-hands in the intact protein are crucial for maintaining its high Ca2+ affinity. The inter EF-hand contacts favor an optimal arrangement of the Ca²⁺ sites in the bound state and promote preformation of the sites in the apo state. Indeed, in the intact protein there is only a very small structural rearrangement upon Ca²⁺ binding.⁴² In the isolated EF-hand fragments, the free energy of Ca²⁺ binding is reduced because a substantial energetic penalty is paid to drive a large conformational change.

We have shown that a series of Ca^{2+} titrations at different peptide concentrations can be used to resolve the coupled Ca^{2+} -binding and dimerization equilibria. The method can be used for other EF-hand fragments as well as for other systems where ligand binding and dimerization are coupled events. We find significant differences between the two EF-hands of calbindin $D_{\rm 9k}$, with a higher dimerization tendency of the less charged EF1 fragment.

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