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Structure of Guanylyl Cyclase Activator Protein 1 (GCAP1) Mutant V77E in a Ca²⁺-free/Mg²⁺-bound Activator State^{*}

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GCAP1, a member of the neuronal calcium sensor subclass of the calmodulin superfamily, confers Ca²⁺-sensitive activation of retinal guanylyl cyclase 1 (RetGC1). We present NMR resonance assignments, residual dipolar coupling data, functional analysis, and a structural model of GCAP1 mutant (GCAP1^{V77E}) in the Ca²⁺-free/Mg²⁺-bound state. NMR chemical shifts and residual dipolar coupling data reveal Ca²⁺-dependent differences for residues 170-174. An NMR-derived model of GCAP1^{V77E} contains Mg²⁺ bound at EF2 and looks similar to Ca^{2+} saturated GCAP1 (root mean square deviations = 2.0 Å). Ca²⁺-dependent structural differences occur in the fourth EFhand (EF4) and adjacent helical region (residues 164-174 called the Ca²⁺ switch helix). Ca²⁺-induced shortening of the Ca²⁺ switch helix changes solvent accessibility of Thr-171 and Leu-174 that affects the domain interface. Although the Ca^{2+} switch helix is not part of the RetGC1 binding site, insertion of an extra Gly residue between Ser-173 and Leu-174 as well as deletion of Arg-172, Ser-173, or Leu-174 all caused a decrease in Ca²⁺ binding affinity and abolished RetGC1 activation. We conclude that Ca²⁺-dependent conformational changes in the Ca²⁺ switch helix are important for activating RetGC1 and provide further support for a Ca²⁺-myristoyl tug mechanism.

Guanylyl cyclase activating proteins $(GCAPs)^2$ belong to the neuronal calcium sensor (NCS) branch of the calmodulin superfamily (1–3) and regulate Ca²⁺-sensitive activity of retinal guanylyl cyclase (RetGC) in rod and cone cells (4–6). Phototransduction in retinal rods and cones is modulated by intracellular Ca²⁺ sensed by GCAPs (7, 8), and defects in Ca²⁺ signaling by GCAPs are linked to retinal diseases (9). GCAP proteins in the Ca²⁺-free/Mg²⁺-bound state activate RetGC (10), whereas Ca²⁺-bound GCAPs inhibit RetGC at high Ca²⁺ levels maintained in the dark (11–13).

The GCAPs (GCAP1 (6), GCAP2 (14), GCAP3 (15), and GCAP4-8 (16) are all ~200-amino acid residue proteins containing a covalently attached N-terminal myristoyl group and four EF-hand motifs (EF1 through EF4; Fig. 1). Mg²⁺ binds to GCAP1 in place of Ca^{2+} when cytosolic Ca^{2+} levels are below 50 nM in light-activated photoreceptor cells (17). This Ca^{2+} free/Mg²⁺-bound GCAP1 is called the activator form because it activates RetGC (10, 18, 19). The x-ray crystal structure of Ca²⁺-bound GCAP1 (20) and NMR structure of GCAP2 (21) showed that the four EF-hands form two semiglobular domains (EF1 and EF2 in the N-domain and EF3 and EF4 in the C-domain); Ca²⁺ is bound at EF2, EF3, and EF4, and the N-terminal myristoyl group in GCAP1 is buried inside the Ca²⁺-bound protein flanked by hydrophobic residues at the N and C termini (see the red residues in Fig. 1). The structure of the physiological activator form of GCAPs (Mg²⁺-bound/Ca²⁺-free state) is currently unknown.

The structure of the Ca²⁺-free/Mg²⁺-bound activator state of GCAP1 has remained elusive, in part because it tends to aggregate under conditions for NMR or x-ray crystallography (22). Here, we present a NMR structural analysis of Ca²⁺-free/ Mg²⁺-bound GCAP1 mutant that has Val-77 replaced by Glu (called GCAP1^{V77E}). The GCAP1^{V77E} mutant retains functional Mg^{2+} and Ca^{2+} binding with intact tertiary structure. However, unlike the dimeric wild type GCAP1, GCAP1^{V77E} is monomeric in solution and remains soluble under NMR conditions. Our NMR analysis indicates that Ca²⁺-free/Mg²⁺bound GCAP1^{V77E} is overall structurally similar to that of Ca^{2+} -saturated GCAP1 (root mean square deviations of 2.0 Å), except that Mg^{2+} is bound at EF2, and the other EF-hands are unoccupied. The largest Ca2+-dependent structural differences in GCAP1 are seen for residues in EF4 and the adjacent helical region (residues 164-174, called Ca²⁺ switch helix). We propose that the Ca^{2+} switch helix may serve as a conduit that relays Ca2+-induced structural changes in EF4 to the RetGC binding site in the N-terminal domain, which provides further support of a Ca^{2+} -myristoyl tug mechanism (23).

Experimental Procedures

Expression and Purification of GCAP1 and Mutants—Mutations were introduced in a bovine GCAP1 coding plasmid using a "splice by overlap extension" approach as previously described (24). Myristoylated GCAP1 and its mutants were produced in *Escherichia coli* strain harboring yeast *N*-myristoyl



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The atomic coordinates and structure factors (code 2NA0) have been deposited in the Protein Data Bank (http://wwpdb.org/). A complete list of NMR assignments for Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} has

A complete list of NMR assignments for Ca^{2+} -free/Mg²⁺-bound GCAP1^{V77E} has been deposited in the Biological Magnetic Resonance Bank (BMRB) (accession no. 26688).

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² The abbreviations used are: GCAP1, guanylyl cyclase activating protein 1; NCS, neuronal calcium sensor; HSQC, heteronuclear single quantum coherence; ITC, isothermal titration calorimetry; RDC, residual dipolar coupling; RetGC, retinal guanylyl cyclase; CaM, calmodulin.



FIGURE 1. Amino acid sequence alignment of bovine GCAP1, recoverin, and NCS-1. Secondary structural elements (α-helices and β-strands) were derived from NMR analyses (22, 37). The four EF-hands (EF1, EF2, EF3 and EF4) are highlighted *green*, *red*, *cyan*, and *yellow*.

transferase and purified to homogeneity using a previously described method (10). The expression and purification of isotopically labeled GCAP1 and mutants were described previously (22, 25). Uniformly ¹⁵N-labeled GCAP1, ¹⁵N-labeled GCAP1^{V77E}, and triple-labeled ¹³C,²H,¹⁵N-labeled GCAP1^{V77E} used in the NMR studies (0.5 mM) were dissolved in 5 mM Tris- d_{11} (pH 7.4), 5 mM CaCl₂, 5 mM MgCl₂, 5 mM dithiothreitol- d_{10} , and 93%/7% H₂O/D₂O.

Mutagenesis—Mutations were introduced in bovine GCAP1 cDNA by "splicing by overlap extension" technique using PCR reactions catalyzed by high-fidelity Phusion Flash polymerase (Finnzymes/Thermo Scientific). The resultant products were ligated into the NcoI/BamHI sites of *pET11d* (Novagen/Calbiochem) vector, sequenced, and transformed into expressing cell lines as described previously in detail (19).

Trp Fluorescence Spectroscopy—The intrinsic Trp fluorescence of GCAP1 and its mutants was recorded in the presence of variable-free Mg^{2+} and Ca^{2+} concentrations as previously described in detail (19, 23).

 Ca^{2+} Binding Stoichiometry—The stoichiometry of Ca²⁺ binding to myristoylated and non-acylated GCAP1 and its mutants was determined using the fluorescent Ca²⁺ indicator dye method previously described in detail (19, 23) using Fluo-4FF (Molecular Probes/Fisher). Free Ca²⁺ in the reaction mixture was calculated using the formula [Ca]_{free} = $K_d \times (F - F_{min})/(100 - F)$, where *F* is the fluorescence intensity of the Ca²⁺ indicator in the assay mixture expressed as a percentage of the fluorescence of the Ca²⁺-saturated indicator (recorded at the end of each experiment in 1 mM [Ca]_{free}), F_{min} is the fluorescence intensity of the Ca²⁺ indicator in the absence of Ca²⁺ and also expressed as a percentage of the fluorescence of the Ca²⁺-saturated indicator, and K_d is a corrected constant of the indicator dye for Ca²⁺ (19). The fluorescence data were fitted by the equation ([Ca]_{bound}/[GCAP]) = $N \times [Ca_{free}]^n ([Ca_{free}]^n + K_d^n)$, where [Ca]_{bound} is the concentration of Ca²⁺ bound to GCAP1 calculated as $[Ca]_{bound} = [Ca]_{total} - [Ca]_{free}$, where *N* is the number of Ca²⁺ ions bound per molecule of GCAP at saturation, K_d is the apparent affinity of GCAP1 for Ca²⁺, and *n* is the Hill coefficient. The data shown are representative from independent experiments producing virtually identical results.

Guanylyl Cyclase Assay—Recombinant human RetGC1 was expressed in HEK293 cells and assayed *in vitro* using $[\alpha^{-32}P]$ GTP as a substrate as previously described in detail (10, 23). Ca²⁺/Mg²⁺/ethylene glycol tetraacetic acid (EGTA) buffers for the assay were prepared and calibrated as previously described (10).

NMR Spectroscopy—Samples for NMR analyses were prepared by dissolving unlabeled, ¹⁵N-labeled, ¹³C, ¹⁵N-labeled, or ¹³C, ²H, ¹⁵N-labeled GCAP1 proteins in 0.5 ml of 90% H₂O, 10% [²H]H₂O containing 10 mM [²H₁₁]Tris (pH 7.4) and either 5 mM MgCl₂ (Mg²⁺-bound) or 5 mM CaCl₂ (Ca²⁺-bound). All NMR experiments were performed at 37 °C on a Bruker Avance 800-MHz spectrometer equipped with a triple resonance cryoprobe and *z* axis gradient. NMR experiments and backbone assignments for Ca²⁺-saturated GCAP1 were described elsewhere (22). Backbone NMR resonance assignments of the Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} activator state (Mg²⁺ bound at EF2) were obtained in this study by the analysis of three-dimensional NMR data as described previously (26). Two-dimensional ¹H, ¹⁵N HSQC with 2048 (¹H) × 256 (¹⁵N) data points, three-dimensional HNCACB with 1024 (¹H) × 64 (¹⁵N) × 120 (¹³C)

TABLE 1	
NMR structural statistics for Mg ²⁺ -bound/Ca ²⁺ -free GCAP1 ^{V77E}	

NMR constraints	GCAP1 ^{V77E}
Distance restraints	
Inter-residue	
Sequential $(i - j = 1)$	500
Medium range $(2 \le i - j \le 4)$	260
Long range $(i - j \ge 5)$	9
Mg ²⁺ -EF-hand	5
Dihedral angle restraints	
φ	117
ψ	117
RDC restraints	41

data points, and three-dimensional HNCO with 2048 (¹H) × 64 (¹⁵N) × 128 (¹³C) data points were all performed on a triplelabeled GCAP1^{V77E} sample using a 800-MHz Bruker NMR spectrometer equipped with a triple resonance cryogenic probe. In addition, three-dimensional HNCOCACB with 1024 (¹H) × 64 (¹⁵N) × 128 (¹³C) data points was performed on the triple-labeled sample using a 600-MHz Bruker NMR spectrometer equipped with a triple resonance cryogenic probe. ¹H, ¹⁵N residual dipolar coupling constants (D_{NH}) were measured with a ¹⁵N-labeled GCAP1^{V77E} (~0.3 mM) containing 12 mg/ml Pf1 phage (Asla Biotech) and using a two-dimensional IPAP (inphase/antiphase) ¹H, ¹⁵N HSQC experiment (27). Spectra were processed using NMRPipe software package (28) and analyzed using SPARKY.

NMR-guided Homology Modeling-An NMR-guided homology model structure of Mg²⁺-bound/Ca²⁺-free GCAP1^{V77E} was generated based on NMR data (chemical shifts, residual dipolar couplings (RDCs) and NOEs) using the Xplor-NIH software suite (29, 30). A template structure for the model calculation was first built using SWISS-MODEL based on the x-ray crystal structure of Ca²⁺-bound GCAP1 (PDB ID 2R2I). The N-terminal myristoyl group was attached to the template structure as it is in the crystal structure (PDB ID 2R2I). The three Ca^{2+} ions were deleted, and a single Mg^{2+} ion was added to the second EF-hand metal binding loop as described by Park et al. (31). The template structure of Mg^{2+} -bound GCAP1^{V77E} then served as input for restrained molecular dynamics (32) followed by refinement (using refine.py) within the Xplor-NIH software suite (30). RDC and NMR restraints (NOEs and dihedral angles) were applied during the simulated annealing step. Dihedral angles were calculated by the Talos+ program (33). For under-assigned secondary structural motifs, theoretical restraints within the initial template structure were used to supplement the experimental restraints. Refinement of the final structure was initiated with high temperature annealing at 1000 K for 10 ps, and cooling from 1000 K to 25 K in 12.5 K steps. The duration of cooling dynamics run at each step was 0.2 ps. A total of 500 structures were obtained, and the 75 lowest energy structures were chosen to generate an energy-minimized average structure. The structural statistics are shown in Table 1.

Isothermal Titration Calorimetry—ITC experiments were performed using a VP-ITC calorimeter (Micro-Cal) at 30 °C, and data were acquired and processed with MicroCal software as described previously (34). Metal-free GCAP1^{V77E} samples were prepared by exchanging protein into buffer containing 15 mM Tris-HCl (pH 7.5), 100 mM NaCl, and 1 mM β -mercaptoethanol. The metal-free GCAP1^{V77E} in the sample cell (50 μ M, 1.5 ml) was titrated with either Ca²⁺ (2 mM) or Mg²⁺ (40 mM) using 40 injections of 5 μ l each.

Results

GCAP1 Mutant (V77E) Binds Functionally to Mg^{2+} and Ca^{2+} —ITC was used to monitor Mg²⁺ and Ca²⁺ binding to GCAP1^{V77E} (Fig. 2). Titration of Mg²⁺ into apoGCAP1^{V77E} produced an endothermic isotherm (Fig. 2A). Mg^{2+} binds to $\text{GCAP1}^{\text{V77E}}$ with an apparent dissociation constant (K_d) of 700 μ M and enthalpy difference (Δ H) of +4.2 kcal/mol. The stoichiometry of Mg²⁺ binding was determined by analyzing ¹H,¹⁵N HSQC NMR spectra of Mg²⁺-bound GCAP1^{V77E}. NMR not only determined the number of Mg²⁺ ions bound per protein but also determined which particular EF-hands are bound to Mg^{2+} (35). The NMR spectrum of Ca^{2+} -free/ Mg^{2+} -bound $GCAP1^{V77E}$ contains one downfield NMR peak at ~10.5 ppm assigned to Gly-69 (conserved glycine in EF2 binding loop), indicating that 1 Mg^{2+} is bound to GCAP1^{V77E} at EF2. (Fig. 2A, *inset*). Thus, GCAP1^{V77E} binds to Mg^{2+} with the same affinity and stoichiometry as wild type GCAP1 (25), which demonstrates that Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} is structurally and functionally intact.

Titration of CaCl₂ into apoGCAP1^{V77E} showed an ITC isotherm (Fig. 2B) with three separate phases that corresponded to Ca²⁺ binding at three EF-hand binding sites (EF2, EF3, and EF4). The Ca²⁺ binding isotherm of GCAP1^{V77E} looked overall similar to that of wild type (25). The highest affinity site in GCAP1^{V77E} ($K_d = 0.08 \ \mu\text{M}$ and $\Delta H = -3 \ \text{kcal/mol}$) exhibited binding energetics that are nearly identical to those of EF3 in wild type (25). Therefore, this highest affinity site in GCAP1^{V77E} was assigned to EF3. The site with intermediate affinity ($K_d = 0.2 \ \mu \text{M}$ and $\Delta H = +1.4 \ \text{kcal/mol}$) was assigned to EF4 based on its similar binding affinity to EF4 in wild type under the conditions of microcalorimetry assay (25). By the process of elimination, the lowest affinity site in GCAP1^{V77E} $(K_d = 10 \ \mu \text{M} \text{ and } \Delta H = -0.5 \text{ kcal/mol})$ was assigned to EF2. The Ca^{2+} binding affinity for EF2 in GCAP1^{V77E} is 10-fold weaker than that of wild type, consistent with the V77E mutation residing in EF2. The lower affinity Ca²⁺ binding to EF2 may also result from a loss of protein dimerization for $GCAP1^{V77E}$ compared with the dimeric wild type protein (22). For example, intermolecular dimer interactions involving EF2 could stabilize Ca²⁺ binding to EF2 in the dimeric wild type protein. Indeed, the ΔH for Ca²⁺ binding to both EF2 and EF4 in GCAP1^{V77E} have opposite signs compared with that of wild type. The opposite sign of ΔH for both EF2 and EF4 in GCAP1^{V77E} suggests that exposed residues in EF2 and EF4 might form Ca²⁺-dependent intermolecular contacts with each other in the dimeric wild type protein that get disabled (or otherwise altered) in the monomeric $GCAP1^{V77E}$ (22).

The stoichiometry of Ca^{2+} binding was determined by analysis of ¹H, ¹⁵N HSQC NMR spectra of Ca^{2+} -bound GCAP1^{V77E} (Fig. 2*B, inset*). Three downfield NMR peaks assigned to Gly-69 (EF2), Gly-105 (EF3), and Gly-149 (EF4) demonstrated that three Ca^{2+} ions are bound per protein at EF2, EF3, and EF4. Thus, GCAP1^{V77E} binds to Ca^{2+} with the same stoichiometry and structure as wild type GCAP1 (17, 36).





FIGURE 2. **GCAP1 mutant GCAP1^{V77E} binds to Mg²⁺ and Ca²⁺ as measured by ITC and NMR (***inset***). ITC binding isotherms recorded at 30 °C are shown for Mg²⁺ binding (***A***) and Ca²⁺ binding (***B***) to GCAP1^{V77E}. Binding isotherms were fit to a sequential model (***solid line***), and fitting parameters are given in "Results." Downfield spectral regions of ¹H,¹⁵N HSQC spectra of Mg²⁺-bound GCAP1^{V77E} (***inset***,** *A***) and Ca²⁺-bound GCAP1^{V77E} (***inset***,** *B***) are also shown. The downfield peak assigned to Gly-69 indicates Mg²⁺ is bound at EF2 (***inset***,** *A***), whereas peaks assigned to Gly-69, Gly-105, and Gly-149 indicate Ca²⁺ is bound at EF2, EF3, and EF4.**

NMR Assignments for Ca^{2+} -free/Mg²⁺-bound GCAP1^{V77E}— NMR spectroscopy was used to demonstrate that GCAP1^{V77E} adopts a native tertiary structure as determined by comparing ¹H, ¹⁵N HSQC spectra of wild type protein (Fig. 3A and Ref 37) to that of GCAP1^{V77E} (Fig. 3, *B* and *C*). The peaks in the 1 H, 15 N HSQC NMR spectra represent main chain and side-chain amide groups that provide a residue-specific fingerprint of the overall protein conformation. The resonance assignments for Ca²⁺-saturated GCAP1^{WT} (37) match quite well with the spectrum of Ca^{2+} -saturated GCAP1^{V77E} (Fig. 3*C*), indicating that GCAP1^{V77E} is structurally intact. The NMR assignments for Ca^{2+} -free/Mg²⁺-bound GCAP1^{V77E} are shown by the labeled peaks in Fig. 3B. The Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} forms a monomer in solution under NMR conditions (22), in contrast to Ca2+-free/Mg2+-bound wild type GCAP1 that tends to aggregate and/or form dimers under NMR conditions (22). The relative peak positions and spectral patterns overall look similar when comparing spectra of wild type (Fig. 3A) to that of GCAP1^{V77E} (Fig. 3B), indicating that Mg^{2+} -bound/ Ca²⁺-free GCAP1^{V77E} retains the same main chain fold compared with wild type. The monomeric state of GCAP1^{V77E} caused much sharper NMR line width s compared with that of dimeric wild type (Fig. 3, A versus B). The sharper NMR peaks observed for Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} allowed ~60% assignment of backbone resonances compared with <25% assignment for Ca²⁺-free wild type and $\sim80\%$ assignment for Ca²⁺-saturated GCAP1 (37). The unassigned residues for GCAP1^{V77E} included the first 20 residues from the N terminus and unstructured regions (Lys-46-Trp-51, Met-74-Asp-108, Ile-119-Met-129, Asp-143-Leu-150, Arg-177-Gln-183). These unassigned residues have weak NMR intensities due to

exchange broadening, suggesting that these residues undergo dynamical motions on the chemical shift time scale. The exchange broadening of residues at the domain interface in Ca^{2+} -free/Mg²⁺-bound GCAP1 (residues Lys-91–Tyr-99) is similar to that seen in Ca^{2+} -free recoverin (38, 39).

Downfield-shifted NMR peaks at ~10.5 ppm for GCAP1^{V77E} are assigned to conserved glycine residues in the EF-hand Ca²⁺ binding loops and are characteristic of Ca²⁺/Mg²⁺-bound EF-hands. In Ca²⁺-saturated GCAP1^{V77E} (Fig. 3C), three downfield peaks assigned to Gly-69 (EF2), Gly-105 (EF3), and Gly-149 (EF4) confirm that Ca²⁺ is bound at EF2, EF3, and EF4 as seen in the crystal structure (20). In Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} (Fig. 3*B*), one downfield peak assigned to Gly-69 (EF3) confirmed that Mg²⁺ is bound at EF2 as seen by fluorescence spectroscopy (10, 19).

NMR-derived Structural Model of Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E}—The sequence-specific NMR assignments above for Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} and the secondary structure based on these assignments is summarized in Fig. 1. An NMR-derived structural model of Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} was calculated using NOE-based distances, NMRderived dihedral angle restraints, and RDC data (Fig. 4) that served as input for restrained molecular dynamics structure calculations (see "Experimental Procedures"). Initial residual dipolar coupling magnitude and rhombicity were calculated by fitting the measured residual dipolar couplings to the calculated structure using the PALES program (40). The RDC-refined structures have a quality Q-factor of 0.28 and an R-factor of 0.995 (Fig. 4C). The overall secondary structure and topology of Mg²⁺-bound GCAP1^{V77E} is similar to that found in the crystal structure of Ca2+-bound GCAP1 (20). The NMR-derived



FIGURE 3. **NMR spectroscopy of GCAP1^{WT}, GCAP1^{V77E}, and mutants (SGL and \DeltaLeu-174).** Two-dimensional (¹H, ¹⁵N HSQC) NMR spectra of ¹⁵N-labeled wild type GCAP1 in the Ca²⁺-free/Mg²⁺-bound state (*A*), GCAP1^{V77E} in the Ca²⁺-free/Mg²⁺-bound state (*B*), GCAP1^{V77E} in the Ca²⁺-bound state (*C*), and GCAP1 mutants (SGL, *black*; Δ Leu-174, *red*) in the Ca²⁺-bound state (*D*). Spectra were obtained at 37 °C. A downfield resonance at ~10.5 ppm for Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} is assigned to a conserved glycine residue (Gly-69) in EF2 and indicates Mg²⁺ is bound at EF2. Downfield resonances (at 10.45, 10.47, and 10.55 ppm) for Ca²⁺-bound GCAP1^{V77E} are assigned to conserved glycine residues (Gly-105, Gly-69, and G149) and indicate that three Ca²⁺ are bound per protein at EF2, EF3, and EF4. Sequence-specific resonance assignments for Ca²⁺-free/Mg²⁺-bound V77E are indicated by the peak labels. Complete NMR assignments were deposited in the BMRB (accession no. 26688).

structural model of Ca^{2+} -free/Mg²⁺-bound GCAP1^{V77E} (PDB ID 2NA0) was validated with PROCHECK: 86% of residues belonged to the most favorable region in the Ramachandran plot.

The NMR-derived structural model of Ca²⁺-free/Mg²⁺bound GCAP1^{V77E} (Fig. 5) contains a total of 11 α -helices and 4 β-strands: α1 (residues 7–14), α2 (18–28), α3 (36–43), α4 (51– 63), $\alpha 5$ (73–83), $\alpha 6$ (90–99), $\alpha 7$ (109–119), $\alpha 8$ (132–143), $\alpha 9$ $(153-161), \alpha 10 (164-174), \alpha 11 (176-183), \beta 1 (33-35), \beta 2$ (70–72), β 3 (106–108), and β 4 (150–152). GCAP1^{V77E} contains two separate domains comprising four EF hands: EF1 (green, residues 18 - 43) and EF2 (red, residues 51 - 83) form the N-terminal domain; EF3 (cyan, residues 90-119) and EF4 (yellow, residues 132-161) form the C-terminal domain. Two C-terminal helices are downstream of EF4 (α 10 and α 11 in Fig. 4). The helix immediately adjacent to EF4 (α 10, *highlighted in orange* in Fig. 5*B*) is one-half turn longer in Ca^{2+} -free/Mg²⁺bound GCAP1 V77E compared with that of Ca²⁺-bound GCAP1 (Fig. 6). The C-terminal helix (α 11) has the same length in both Ca²⁺-free and Ca²⁺-bound GCAP1 and makes contacts with the N-terminal myristoyl group (Fig. 5).

GCAP1^{V77E} contains Mg²⁺ bound at EF2 (*blue sphere* in Fig. 5) as evidenced by characteristic Mg²⁺-dependent amide

chemical shift changes assigned to Gly-69 in EF2 (Fig. 2*A*, *inset*). The geometry of the coordinate covalent bonds formed between chelating amino acid residues in GCAP1^{V77E} and the bound Mg^{2+} could not be observed directly in our NMR study. Instead, the stereochemical geometry and chelation of Mg^{2+} bound at EF2 was modeled with structural constraints derived from the x-ray crystal structure of Mg^{2+} -bound CaM (41), which closely resembles the Mg^{2+} binding site geometry conserved in other EF-hand proteins such as CaBP1 (35) and CaBP4 (31). GCAP1 residues at the 1, 3, and 5 positions of the EF-hand loop in EF2 were selected to chelate the bound Mg^{2+} (see *D64* and *D68* in Fig. 5*A*). Replacement of these two residues disrupted Mg^{2+} binding in our previous biochemical studies (19).

The four EF-hands of GCAP1^{V77E} with one Mg²⁺ bound at EF2 (and no metal bound at EF1, EF3, and EF4) each adopt interhelical angles that are similar to those observed in the crystal structure of Ca²⁺-bound GCAP1 (Table 2). For the Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} structure, the interhelical angles are 132° (EF1), 114° (EF2), 103° (EF3), and 106° (EF4). Therefore, the three functional EF-hands in GCAP1 (EF2, EF3, and EF4) each adopt a somewhat open conformation in the Ca²⁺-free state, and Ca²⁺ binding at these sites in





FIGURE 4. **RDC structural analysis of GCAP1**^{V77E}. ¹H,¹⁵N IPAP (inphase/antiphase)-HSQC spectra of Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} in the absence (*A*) and presence (*B*) of 12 mg/ml Pf1 phage. Spectral splittings for the isotropic condition (J_{NH}) *versus* the anisotropic condition (J_{NH}) + D_{NH}) are marked by *vertical lines* and were used to calculate RDCs as described under "Experimental Procedures." *C*, RDCs calculated from the structure of Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} in Fig. 5 are plotted *versus* the RDCs measured in Fig. 4*B* and show good agreement (Q-factor = 0.28 and an R-factor = 0.95 (40)).

GCAP1 cause only a slight change in interhelical angle. In essence, the three functional EF-hands in GCAP1 adopt a "pre-formed" open conformation in the Ca²⁺-free state akin to that of calbindin D_{9k} (42) and other Ca²⁺ buffer proteins (43). As a result, the Ca²⁺-binding free energy for GCAP1 is NOT coupled to an unfavorable conformational change, which may explain the very high nanomolar Ca²⁺ binding affinity for GCAP1 (25). By stark contrast, myristoylated recoverin (that undergoes a Ca²⁺-myristoyl switch (44)) and CaM (that undergoes an open-to-closed transition (43)) both bind Ca²⁺ with ~100-fold lower affinity compared with myristoylated GCAP1 (25).

 Ca^{2+} -induced Conformational Changes in GCAP1—Chemical shift differences in GCAP1 versus Ca²⁺ were plotted as a function of residue number and reveal the location of Ca²⁺-dependent structural changes (Fig. 6A). Detectable chemical shift differences are seen for residues in EF1 (residues 26, 27, and 33) that are implicated in RetGC1 binding (36). Somewhat larger chemical shift changes are seen in EF2 (residues 55, 62, and 68), which represent residues at the domain interface and suggest Ca²⁺-dependent structural contacts between EF2 and EF3 like that seen previously for recoverin (38, 45). The largest chemical shift differences (highlighted in *red* in Fig. 6B) are observed in EF4 (residues 140, 151–152, and 160) and helix α 10 (residues



FIGURE 5. **NMR-derived structure of Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} (PDB ID 2NA0).** The main chain structure of Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} (A) and the same view rotated by 180 degrees (B) show four EF-hands (colored as in Fig. 1) packed in a globular arrangement very similar to what is seen for Ca²⁺-bound GCAP1 (20). The secondary structural elements are labeled as defined in Fig. 1. The Ca²⁺ switch helix (α 10) is highlighted in *orange*, bound Mg²⁺ is in *blue*, and the N-terminal myristoyl group is in *magenta*.

168, 170, and 171), called the Ca^{2+} switch helix. Thus, Ca^{2+} induced structural changes in EF4 appear coupled to structural changes in the adjacent Ca²⁺ switch helix. An overlay of GCAP1 structures in the Ca²⁺-free versus Ca²⁺-bound states shows overall main chain root mean square deviations of 2.0 Å. The most striking difference is seen in the Ca^{2+} switch helix (orange in Figs. 5B and 6C), which is one-half-turn longer in the Ca^{2+} -free state. An expanded view of the Ca^{2+} switch helix (Fig. 6C) reveals two residues (Thr-171 and Leu-174) that are most affected by Ca^{2+} . Thr-171 is exposed in the Ca^{2+} -free structure, whereas it becomes buried and makes contact with Leu-92 in the Ca²⁺-bound structure. Conversely, Leu-174 is buried and makes contact with Leu-92 in the Ca²⁺-free structure, but it switches to a solvent-exposed environment in the Ca²⁺-bound structure. The Ca²⁺-dependent contacts formed by both Thr-171 and Leu-174 may be important for switching

GCAP1 from the Ca²⁺-free activator to the Ca²⁺-bound inhibitor states.

Mutations in the Ca²⁺ Switch Helix Affect Metal Binding and Cyclase Activation—The calcium-myristoyl tug hypothesis (23) proposes a structural link between the exiting helix of EF4 and residues that contact the myristoyl group in the N-domain. Ca²⁺-induced structural changes in EF4 exert a "tug" on downstream C-terminal residues that are in contact with the myristoyl group. These Ca²⁺-dependent contacts to the myristate are then relayed to the target binding site in the N-domain (36). We hypothesized that the Ca²⁺-induced shortening of the Ca²⁺ switch helix observed in our structure (Fig. 6) may transmit the tug action that connects Ca²⁺ binding at EF4 with target activation in the N-domain. Therefore, we tested if a change in the length of the Ca²⁺ switch helix by mutagenesis could affect the regulatory properties of GCAP1. We constructed a





FIGURE 6. **Ca²⁺-induced conformational changes in GCAP1.** Shown in Ca²⁺-dependent amide chemical shift difference (Ca²⁺-free minus Ca²⁺-bound) plotted *versus* residue number (*A*) and chemical shift difference mapped onto the main chain structure (*B*). Residues Thr-171 and Leu-174 exhibited the largest Ca²⁺-induced chemical shift differences. Residues (Thr-62, Phe-140, Leu-151, Val-160, Leu-170, Thr-171, Leu-174) with a chemical shift difference higher than 0.8 are colored *red*. Residues (Ala-52, Tyr-55, Asp-68, Ile-115, Ala-118, Ser-152, Glu-158, Gln-161, Asp-168) with a chemical shift difference between 0.5 and 0.8 are colored *magenta*. Residues with a chemical shift difference < 0.5 are colored light blue. C, close-up view of the Ca²⁺ switch helix (*a*10, *orange*) that is elongated by one turn in Ca²⁺-free/Mg²⁺-bound GCAP1^{VYTE} (*left*) compared with Ca²⁺-bound GCAP1^{WT} (*right*). The angle between EF2 exiting helix (*red*) and EF3 entering helix (*cyan*) at the domain interface increased slightly (*dotted line*) due to Ca²⁺-dependent interactions with the Ca²⁺ switch helix.

series of mutants in which the length of the helix was altered either by insertion or deletion of a single amino acid: a Gly was inserted between Ser-173 and Leu-174 (SGL mutant, Fig. 7), and single amino acids were deleted: Arg-172 (Δ Arg-172), Ser-173 (Δ Ser-173), or Leu-174 (Δ Leu-174) (Fig. 8). NMR HSQC spectra demonstrate that these mutations do not alter the conformation of α 10 and these mutants are structurally intact (Fig. 3*D*). We found that each of these mutants (which changes the length of the Ca²⁺ switch helix) profoundly affected both the metal binding affinity of GCAP1 and its ability to activate RetGC. The Ca²⁺ binding stoichiometry in the SGL mutant remained three per GCAP1 molecule, but the apparent affinity decreased slightly with higher cooperativity compared with that of wild type (Fig. 6A). Metal-dependent changes in the intrinsic Trp fluorescence of the SGL mutant looked quite different from that of wild type (Fig. 7, B-D). The Ca²⁺-induced decrease in fluorescence (in the absence of Mg²⁺) occurred at higher Ca²⁺ levels in SGL compared with wild type (Fig. 7*B*). The presence of 10 mM Mg²⁺ (that normally causes a quite striking Ca²⁺-induced increase in Trp fluorescence for wild type) resulted in much smaller Ca²⁺-dependent change in Trp fluorescence for SGL (Fig. 7*C*). Thus, the transition from the Mg²⁺-bound activator state to the Ca²⁺-bound inhibitor state in the presence of 10 mM Mg²⁺ (19) was more difficult to mon-

TABLE 2 Interhelical angles of the EF-hands in GCAP1 and CaM

Residues in the helices are as shown in the footnotes.

	Interhelical angles				
Helix pairs	ApoCaM ^a	Ca ²⁺ -bound CaM ^b	Mg ²⁺ -bound GCAP1 ^c	Ca ²⁺ -bound GCAP1 ^c	
	Degree				
EF1 $(\alpha_1 - \alpha_2)$	130.9	103.8	132.1	132.7	
EF2 $(\alpha_3 - \alpha_4)$	130.8	101.0	113.9	112.1	
EF3 $(\alpha_5 - \alpha_6)$	139.5	101.0	102.9	103.1	
EF4 ($\alpha_7 - \alpha_8$)	126.0	101.0	106.1	104.0	

^a ApoCaM (PDB accession code 1DMO): (α1) 6–18, (α2) 29–38, (α3) 45–55, (α4) 65–75, (α5) 82–90, (α6) 103–112, (α7)118–127, (α8) 137–143.
 ^b Ca²⁺-bound CaM (PDB accession code 1J7P): (α1) 6–19, (α2) 29–38, (α3) 45–55, (α4) 65–75, (α5) 83–92, (α6) 102–111, (α7)118–128, (α8) 138–145.
 ^c GCAP1: (α₁) 18–27, (α₂) 36–43, (α₃) 51–63, (α₄) 73–82, (α₅) 90–99, (α₆) 109–119, (α₇) 132–143, (α₈) 153–161.

itor in SGL compared with wild type (Fig. 7*C*). In addition, the Trp fluorescence of the SGL mutant titrated as a function of Mg^{2+} revealed a nearly 10-fold lower Mg^{2+} binding affinity compared with that of wild type (Fig. 7*D*). Because Mg^{2+} binding to EF2 is essential for converting GCAP1 into its activated state in the absence of Ca^{2+} (10, 19), we tested if RetGC1 activation was also affected by this mutation. Indeed, the SGL mutant failed to activate RetGC1 at physiological concentrations of Mg^{2+} and when the Ca^{2+} concentration was reduced to levels typical of light-exposed photoreceptors (Fig. 7*D*).

Making the Ca²⁺ switch helix shorter by a single residue produced a similar effect regardless of which amino acid was deleted, Arg-172, Ser-173, or Leu-174 (Fig. 8). In all three deletion mutants, the Ca²⁺-induced decrease in Trp fluorescence looked similar to that of wild type (Fig. 8, A-C, red symbols), except for a slight 2-fold shift toward higher Ca²⁺ concentrations (Fig. 8D). As seen above for SGL, the presence of 10 mM Mg²⁺ caused a much smaller change in Ca²⁺-dependent Trp fluorescence for each of the deletion mutants, thus making it difficult to monitor the transition from the Mg²⁺-bound activator state to the Ca^{2+} -bound inhibitor state (Fig. 8, A-C, blue symbols) in contrast to the much larger Ca²⁺-induced fluorescence change seen for wild type (Fig. 7B). As seen above for SGL, the Trp fluorescence of the deletion mutants titrated as a function of Mg²⁺ revealed a nearly 10-fold lower Mg²⁺ binding affinity compared with that of wild type (Fig. 8E). Finally, the deletion mutants all failed to activate RetGC1 at physiological concentrations of Mg^{2+} , consistent with their lack of Mg^{2+} binding (Fig. 8F).

Discussion

NCS proteins like recoverin and NCS-1 undergo large Ca²⁺induced conformational changes that cause extrusion of the N-terminal myristoyl group, termed Ca²⁺-myristoyl switch (38, 46–48). Surprisingly, the GCAP proteins do not possess a Ca²⁺-myristoyl switch (17, 49, 50), and the N-terminal myristoyl group remains buried inside the protein in both Ca²⁺-free and Ca²⁺-bound GCAPs (17, 51). Although NMR and crystal structures are known for Ca²⁺-bound GCAPs (20–22, 52), before this study little was known structurally about the Mg²⁺bound/Ca²⁺-free activator state of GCAPs and how Ca²⁺-induced conformational changes in GCAPs control cyclase activation.

Structure of Ca²⁺-free/Mg²⁺-bound GCAP1

In this study we present NMR assignments (Fig. 3*B*), NOEs (Table 1), RDCs (Fig. 4), and mutagenesis data (Fig. 7) to probe Ca²⁺-dependent conformational changes between the Ca²⁺saturated inhibitory state versus the Ca²⁺-free/Mg²⁺-bound GCAP1^{V77E} (Fig. 6). Overall, the NMR chemical shift differences between Ca^{2+} -saturated wild type GCAP1 and Ca^{2+} -free/Mg²⁺-bound GCAP1^{V77E} are relatively small (Fig. 6A), consistent with an overall similar main chain conformation for the two states (root mean square deviations = 2.0 Å). This contrasts with the large Ca²⁺-induced conformational changes seen for other EF-hand proteins, including recoverin (38), NCS-1 (48), CaM (53, 54), and troponin-C (55). The large Ca²⁺-induced conformational changes in recoverin and CaM both drive the exposure of hydrophobic residues, which consumes Ca²⁺ binding free energy and accounts for their low binding affinity. The Ca²⁺-free EF-hands in GCAP1 adopt a preformed open conformation, like what is seen for calbindin D9k (42) and other Ca^{2+} buffer proteins (43). The lack of any large scale Ca²⁺-induced protein conformational change in GCAP1, therefore, may explain why the GCAPs bind Ca^{2+} with at least 100-fold higher affinity compared with that of recoverin (44) and CaM.

The most apparent Ca²⁺-induced structural changes in GCAP1 are localized in EF4 and the adjacent Ca²⁺ switch helix (α 10). Ca²⁺ binding to EF4 is essential for switching GCAP1 between activator and inhibitor states (10, 19). Upon Ca²⁺ binding to EF4, its interhelical angle decreases by 2° (Table 2) along with a slight lengthening of the exiting helix (α 9) that in turn exerts a force (tug) on the adjacent Ca²⁺ switch helix (α 10). As a result, the Ca²⁺ switch helix is one half-turn longer in the Ca^{2+} -free structure compared with the Ca^{2+} -bound structure (Fig. 6, *B* and *C*). The Ca^{2+} -induced shortening of the Ca²⁺ switch helix alters the disposition of key residues (Thr-171 and Leu-174) that form Ca²⁺-dependent contacts with EF3 (Leu-92 and Tyr-95). These Ca²⁺-dependent contacts alter the packing angle between helices $\alpha 5$ and $\alpha 6$ at the domain interface (Fig. 6*C*) that resembles the Ca^{2+} -induced domain swiveling in recoverin (38). The Ca^{2+} -dependent structural changes in the Ca²⁺ switch helix also exert a force on the C-terminal helix (α 11) that alters its contact with the myristoyl group. We propose that this Ca²⁺-dependent perturbation of the myristoyl group could alter its contact with EF1 and EF2 and thereby affect the accessibility of GCAP1 residues (Met-26, Tyr-37, Val-77, and Ala-78) in the RetGC1 binding site (36). In essence, the Ca²⁺-induced structural changes in EF4 are relayed to the cyclase binding site via the Ca²⁺ switch helix. This relay scheme is consistent with a previously proposed mechanism called the Ca^{2+} -myristoyl tug (23).

We further find here that even though the Ca²⁺ switch helix (α 10) is not an immediate part of the RetGC binding interface (36), the length of α 10 critically affects proper binding of Ca²⁺ (23) and Mg²⁺ (Figs. 7 and 8). This effect on Mg²⁺ binding (Figs. 7 and 8) is especially important for the RetGC1 binding site in GCAP1, which includes the N-domain and requires Mg²⁺ binding in EF2 to interact with the cyclase at low Ca²⁺ levels that occur in light-adapted photoreceptors (19, 36, 56). Evidently, the length of the Ca²⁺ switch helix (α 10) in wild type GCAP1 has evolved to optimize the proper tug action between





FIGURE 7. Single amino acid residue insertion in the Ca²⁺ switch helix affects metal sensor properties of GCAP1. A, Ca²⁺ binding isotherm for wild type (*black*, \oplus) and SGL (*red*, \oplus) GCAP1. Ca²⁺ binding was assayed using titration of 20 μ M GCAP1 in the presence of Fluo4FF as described under "Experimental Procedures." *B–D*, tryptophan fluorescence titrations for monitoring metal-dependent conformational change in wild type (*black*, \oplus) and SGL (*red*, \oplus) GCAP1 caused by Ca²⁺ (*B* and C) or Mg²⁺ (*D*) binding. *B* and *C*, Ca²⁺ titration in the absence (*B*) or in the presence (*C*) of 10 mM Mg²⁺. *AU*, absorbance units. *D*, Mg²⁺ titration. *E*, RetGC1 activation *in vitro* by wild type (*black*, \oplus) and SGL (*red*, \oplus) GCAP1 in the presence of 1 mM Mg²⁺ was assayed as described under "Experimental Procedures."

the N- and C-domains, which tunes the Ca^{2+} and especially Mg^{2+} binding affinity into the physiological range.

GCAP1 residues in EF2 and EF3 located at the domain interface (Val-77–Trp-94, see Figs. 1 and 6) exhibit exchange-broadened NMR resonances and are conformationally dynamic. The dynamical nature of these residues is consistent with Ca^{2+} induced conformational changes in GCAP1 at the domain interface that were observed previously (13, 57). The corresponding residues in recoverin (Fig. 1) also exhibited broad NMR resonances, and ¹⁵N NMR relaxation dispersion studies revealed that these residues at the domain interface exhibit backbone dynamics on the millisecond timescale (45). Ca^{2+} induced rearrangement of residues at the domain interface in recoverin gives rise to a 45-degree swiveling of the two domains (38). Our structural model of Mg^{2+} -bound/ Ca^{2+} -free GCAP1^{V77E} suggests a related but smaller Ca²⁺-induced rearrangement at the domain interface (Fig. 6). Residues Tyr-55, Thr-62, and Ala-118 at the domain interface exhibit large Ca²⁺-induced chemical shift differences (Fig. 6A). In addition, residues in EF3 (Leu-92 and Tyr-95) make Ca²⁺-dependent contacts with Thr-171 and Leu-174 in the Ca²⁺ switch helix (Fig. 6C). These Ca²⁺-dependent contacts to EF3 cause a change in packing angle between EF2 and EF3 at the domain interface (see the *dotted line* in Fig. 6C) that is somewhat reminiscent of the Ca²⁺-dependent domain swiveling observed for recoverin (38, 45).

NMR relaxation data and size-exclusion chromatography analysis previously showed that GCAP1 is dimeric in solution at high micromolar protein concentration (22). A GCAP1 homolog, GCAP2, undergoes a Ca^{2+} -sensitive dimerization at low



FIGURE 8. Effect of a single amino acid residue deletion in Ca²⁺ switch helix on metal sensor properties of GCAP1. A–C, tryptophan fluorescence titrations for monitoring Ca²⁺-dependent conformational change in Δ Arg-172 (**□**) (A), Δ Ser-173 (**△**) (B), or Δ Leu-174 (**♦**) (C) GCAP1 in the absence (*red symbols*) or in the presence (*blue symbols*) of 10 mm Mg²⁺. AU, absorbance units. D and *E*, comparison of the Ca²⁺-dependent (*D*) or Mg²⁺-dependent (*E*) Trp fluorescence change in the wild type (*black*, **●**) Δ Arg-172 (**□**) (A), Δ Ser-173 (**△**) (B), or Δ Leu-174 (**♦**), does dependent (*D*) or Mg²⁺-dependent (*E*) Trp fluorescence change in the wild type (*black*, **●**) Δ Arg-172 (**□**) (A), Δ Ser-173 (**△**) (B), or Δ Leu-174 (**♦**); no Mg²⁺ added in *D*. *F*, does dependence of RetGC1 activation *in vitro* by wild type (*black*, **●**) Δ Arg-172 (*red*, **△**), or Δ Leu-174 (*red* **◊**) GCAP1 in the presence of 6 mm free Mg²⁺ and 2 mm EGTA; the data points were fitted using Synergy Kaleidagraph 4 software assuming a standard Michaelis hyperbolic function.

micromolar protein concentrations, which originally suggested that reversible dimerization may control formation and activation of RetGC:GCAP in a 2:2 complex (58, 59). Although dimerization of GCAP1 observed by Lim *et al.* (22) does not appear to be Ca^{2+} -sensitive, it is possible that GCAP1 dimerization might promote a functional interaction within a RetGC1 dimer on the disk membrane (60, 61). The binding stoichiometry of the GCAP1:RetGC1 complex has been estimated to be equimolar (62), consistent with a 2:2 complex that contains a RetGC1 dimer bound to a GCAP1 dimer. Previously we showed that the V77E mutation in GCAP1 eliminated GCAP1 dimerization and abolished its ability to bind RetGC but did not block its ability to undergo the functional transition between Ca^{2+} -bound and Mg^{2+} -bound states (22, 36). If GCAP1 dimerization is important for RetGC activation, allosteric changes in the GCAP1 dimer quaternary structure may control cyclase activation. In this scenario a small Ca^{2+} -induced change in tertiary structure could lead to a much larger change in quaternary structure akin to the O₂-dependent conformational changes in hemoglobin (63). Hydrophobic residues at the GCAP1 domain interface (Val-77, Ala-78, Leu-82, Trp-94) are solvent-exposed and might mediate specific contacts that control RetGC1 binding and GCAP1 dimerization alike. However, it cannot be excluded that these exposed GCAP1 res-



idues may create an artificial dimer at high concentrations and in the absence of the target enzyme. Future studies are needed to further probe the dimeric structure of GCAP1 and determine whether Ca^{2+} -induced changes in quaternary structure might control its activation of RetGC1.

Author Contributions—J. B. A. directed the overall project and wrote the paper. S. L. performed NMR and ITC experiments, analyzed the NMR and ITC data, performed structure calculations, and helped write the manuscript. E. V. O. and A. M. D. constructed the mutants. I. V. P. assayed Trp fluorescence, Ca^{2+} binding stoichiometry, and RETGC activity. I. V. P. and A. M. D. analyzed the mutagenesis data and participated in writing the manuscript.

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