Crystal Structures of the GCaMP Calcium Sensor Reveal the Mechanism of Fluorescence Signal Change and Aid Rational Design*S

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The genetically encoded calcium indicator GCaMP2 shows promise for neural network activity imaging, but is currently limited by low signal-to-noise ratio. We describe x-ray crystal structures as well as solution biophysical and spectroscopic characterization of GCaMP2 in the calcium-free dark state, and in two calcium-bound bright states: a monomeric form that dominates at intracellular concentrations observed during imaging experiments and an unexpected domain-swapped dimer with decreased fluorescence. This series of structures provides insight into the mechanism of Ca2+-induced fluorescence change. Upon calcium binding, the calmodulin (CaM) domain wraps around the M13 peptide, creating a new domain interface between CaM and the circularly permuted enhanced green fluorescent protein domain. Residues from CaM alter the chemical environment of the circularly permuted enhanced green fluorescent protein chromophore and, together with flexible inter-domain linkers, block solvent access to the chromophore. Guided by the crystal structures, we engineered a series of GCaMP2 point mutants to probe the mechanism of GCaMP2 function and characterized one mutant with significantly improved signal-to-noise. The mutation is located at a domain interface and its effect on sensor function could not have been predicted in the absence of structural data.

The intracellular concentration of free calcium ions, $[Ca^{2+}]$, is a universal second messenger in excitable cells such as neurons and muscle cells (1). In neurons, action potential firing triggers Ca2+ influx through voltage-gated calcium channels

(2), and synaptic input results in Ca²⁺ influx through neurotransmitter receptors (3). The timing of spikes and patterns of synaptic input can be inferred by observing intracellular [Ca²⁺] (4), allowing researchers to monitor the activity of individual neurons and neural populations by non-invasive imaging techniques, using calcium-sensitive dyes and proteins.

Genetically encoded calcium indicator proteins (also known as fluorescent calcium indicator proteins) offer the possibility of long-term in vivo imaging of neural network activity, with cell-type- and subcellular localization-specific targeting not easily possible with synthetic small molecule calcium indicators (5). In the decade since their first publication, several formats have been explored for genetically encoded calcium indicator construction, including both two-fluorescent protein (FP)² sensors with a fluorescence resonance energy transfer signal change and single-FP intensity-modulated sensors (6). Genetically encoded calcium indicator-enabled calcium imaging has been used to detect single action potentials in awake mice (7, 8). GCaMP-derived single-FP sensors (9) have facilitated in vivo imaging of odor-evoked neural activity in Drosophila (10) and Caenorhabditis (11), visual-evoked tectal responses in zebrafish (12), and cardiac output in mouse (13).

The GCaMP sensor is composed of a circularly permuted enhanced GFP (cpEGFP) moiety attached to the calcium-binding protein calmodulin (CaM) and the CaM-binding peptide M13pep (from myosin light chain kinase) (Fig. 1A). The first GCaMP sensor was extremely dim and folded poorly at 37 °C, limiting its utility for imaging applications. Incremental rounds of improvement by grafting of GFP-stabilizing mutations and random mutagenesis resulted in GCaMP2 (13–15), with significantly improved folding and fluorescence characteristics. This calcium indicator was extensively characterized in acute mouse cortical brain slice, with the conclusion that although useful, GCaMP2 suffers from low baseline fluorescence and cannot reproducibly detect single action potentials due to poor signalto-noise (6, 16).

² The abbreviations used are: FP, fluorescent protein; GFP, green fluorescent protein; EGFP, enhanced GFP; cpEGFP, circularly permuted EGFP; CaM, calmodulin; M13pep, myosin light-chain kinase target peptide; SEC, size exclusion chromatography; BisTris, 2-[bis(2-hydroxyethyl)amino]-2-(hydroxymethyl)propane-1,3-diol; MOPS, 4-morpholinepropanesulfonic acid; PDB, Protein Data Bank.



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supplemental Figs. S1-S10 and Tables S1-S3.

The atomic coordinates and structure factors (codes 3EK7, 3EK4, 3EKJ, 3EKH, and 3EK8) have been deposited in the Protein Data Bank, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (http:// www.rcsb.org/).

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EXPERIMENTAL PROCEDURES

Protein Expression and Purification—The pRSETa vector containing GCaMP2 was a kind gift of Karel Svoboda (Janelia Farm Research Campus, Howard Hughes Medical Institute). GCaMP2 was expressed and purified as described previously (17). Briefly, BL21(DE3) cells containing pRSETa harboring gcamp2 or gcamp2 mutants were grown in ZYM-5052 medium (18) for 48 h at 25 °C with shaking at 200 rpm. After centrifugation, cell lysis, and clarification, proteins were purified from the cell-free extract by nickel-affinity chromatography. Protein purity over 95% was confirmed by SDS-PAGE analysis. Proteins were dialyzed into 20 mm Tris-HCl, 100 mm NaCl, 2 mm CaCl₂, pH 8.0, and concentrated. Calcium-free samples were prepared identically, except the buffer contained 5 mm EGTA instead of CaCl₂.

Crystallization and Data Collection—All GCaMP crystallization was carried out at 20 °C. All GCaMP protein samples for crystallization were in 20 mm Tris, 100 mm NaCl, 2 mm CaCl₂, pH 8.0, except for the 8EF-apo mutant where the same buffer with 5 mm EGTA substituted for CaCl2 was used. All crystals used for data collection were grown using the hanging-drop vapor diffusion method in 24-well VDX plates. Crystallization of Ca²⁺-saturated dimeric GCaMP2 was described previously (17). The calcium-saturated K378W (at 5.6 mg/ml) and G87R (at 1.5 mg/ml) mutants crystallized ~4 days after mixing with a precipitant solution consisting of 0.1 M magnesium formate dihydrate and 15% polyethylene glycol 3,350 using drop ratios of 2 μ l of protein to 2 μ l of precipitant for K378W and 1.5–2.5 μl for G87R. Ca²⁺-saturated monomeric GCaMP2 was crystallized identically to the K378W and G87R mutants using a drop ratio of 2 μ l to 2 μ l except that the drops were microseeded by streak seeding from K378W crystals immediately following setup. These crystals required more than 4 weeks to grow and had a distinct morphology. 8EF-apo GCaMP2 was crystallized after 1 week by mixing 2 µl of protein solution (9.5 mg/ml) with 2 μl of a precipitant solution consisting of 0.2 M lithium sulfate monohydrate, 0.1 M BisTris, pH 5.5, and 25% polyethylene gly-

All crystals were cryoprotected for data collection by quickly (<10 s) soaking in the precipitant solution supplemented with 20% glycerol and then mounted in a nitrogen gas stream at 100 K or plunged into liquid nitrogen for storage and transport to synchrotron beamlines. All data were collected at 100 K in a $\rm N_2$ gas stream. X-ray diffraction data for the G87R mutant was collected in-house on a Rigaku RU-H3R rotating copper anode x-ray generator, equipped with a Saturn 92 CCD detector and X-stream 2000 low-temperature system. Data for $\rm Ca^{2+}$ -saturated monomeric GCaMP2 was collected at the Advanced Light Source, beamline 8.2.2. Diffraction data from crystals of $\rm Ca^{2+}$ -dimer, K378W, and 8EF-apo GCaMP2 were collected at the Advance Photon Source, beamline 31-ID.

X-ray diffraction data for G87R were integrated and scaled using d*TREK (19) from within the CrystalClear software package (Rigaku/Molecular Structure Corporation, Woodlands, TX). Data for ${\rm Ca}^{2+}$ -saturated monomeric GCaMP2 were integrated and scaled in HKL2000 (20). Data from crystals of ${\rm Ca}^{2+}$ -

dimer, K378W, and 8EF-apo GCaMP2 were processed using Mosflm (21) and Scala (22).

Structure Solution, Model Building, and Refinement-All GCaMP2 structures were solved by molecular replacement using the program Phaser (23). The Ca²⁺-saturated dimer structure was solved as described previously (17) using the published coordinates of GFP (Protein Data Bank (PDB) entry 1EMA) and the coordinates of M13-bound calmodulin (PDB entry 1CDL) as search models. The G87R Ca2+-bound monomer mutant structure was solved by searching sequentially using the cpEGFP domain and CaM-M13 domains from the refined Ca²⁺-dimer structure and data between 29.3- and 2.8-Å resolution. Clear solutions were obtained in space group P4₁2₁2 with translation function Z-scores of 43.1 and 21.3, respectively, for the two domains. Strong positive peaks in the difference map at the expected positions of the calcium ions in CaM (which were omitted from the MR model) indicated the correctness of the solutions. The K378W mutant crystals were isomorphous with those of the G87R mutant and the G87R model was used directly for rigid-body refinement against data from K378W crystals. The Ca²⁺-saturated monomeric GCaMP2 structure was solved using the refined K378W coordinates as a search model. A clear solution was obtained in space group $P2_12_12$ with a translation function Z-score of 39.2 using data between 45.4- and 2.65-Å resolution. The 8EF-apo GCaMP2 calcium-free mutant structure was solved by searching for the cpEGFP domain from the Ca²⁺-dimer structure. A clear solution was obtained in space group C2 with a translation function *Z*-score of 18.9 using data between 31.9- and 2.8-Å resolution. Subsequently searching for the calcium-free N-terminal or C-terminal lobes of CaM (PDB code 1CFD (24)) did not reveal any clear solutions. Some positive difference density was present in the electron density maps calculated using the cpEGFP domain solution that suggested the position of the N-terminal lobe of CaM, which was placed manually into density and refined. The correctness of this CaM N-terminal lobe placement was indicated by additional positive difference density for the linker connecting cpEGFP and the CaM N-terminal lobe, which was subsequently built.

All models were improved by iterative cycles of model building in Coot (25) and positional refinement in REFMAC (26). Final GCaMP2 models have reasonable *R*-factors and model geometries, as illustrated in Table 1. A portion of the electron density map for each structure is provided in supplemental Fig. S1.

Size Exclusion Chromatography (SEC)—All SEC was carried out using a Superdex 200 10/300 GL column (GE Healthcare) at a flow rate of 0.5 ml min $^{-1}$ in 20 mM Tris, 100 mM NaCl, 2 mM CaCl $_2$, pH 8.0, for calcium-saturated samples or with 5 mM EGTA in place of CaCl $_2$ for calcium-free samples. Molecular weights were estimated by comparison with elution volumes of standard proteins (Bio-Rad).

Sedimentation Velocity Analytical Ultracentrifugation—Analytical ultracentrifugation of GCaMP2 samples was carried out in a Beckman XL-I analytical ultracentrifuge (Beckman Coulter, Fullerton, CA) within the Biophysics Instrumentation Facility at the Massachusetts Institute of Technology. Absorbance scans at 280 nm were collected on calcium-free (24 μ M)



and calcium-saturated (28 μm) GCaMP2 samples in two-sector cells within a four-hole AnTi-60 rotor at 42,000 rpm. Data were collected at 20 °C in the same buffers as the SEC experiments. Absorbance scans were modeled using a continuous c(s) distribution within Sedfit (27), correcting for buffer density and viscosity and using a partial specific volume of $0.7300 \text{ cm}^3 \text{ g}^{-1}$. Molecular weight of observed species (Fig. 2B) was estimated based on the best-fit frictional ratio as determined by Sedfit for each sample.

Two-photon Laser Scanning Microscopy of Neurons Expressing GCaMP2-To measure intracellular [GCaMP2], acute brain slices containing neurons expressing GCaMP2 were prepared and imaged as previously described (6, 16). Purified GCaMP2 was diluted into pipette internal solution supplemented with 1 mm K₂EGTA at 0.1, 1, and 10 µm concentrations. Each solution was drawn into a thin glass capillary (ID = 0.02mm, Vitrocom number RT5002). Their fluorescence intensities were measured under two-photon excitation with identical parameters (910 nm excitation) to neuron imaging. 0.1 µM GCaMP2 was not bright enough to significantly exceed PMT dark current at laser powers used for neuronal imaging.

Intracellular GCaMP2 concentration in neurons with robust fluorescent responses to action potential firing was estimated by a linear extrapolation from the purified 10 μM GCaMP2 fluorescence intensity. Intracellular GCaMP2 was assumed to be in the apo state (6).

Generation and Screening of GCaMP2 Mutants—Mutants of GCaMP2 were prepared by site-directed mutagenesis (see supplemental Tables S1-S3) and confirmed by sequencing. Preliminary screening for variants with altered oligomerization equilibria (supplemental Table S1) was performed by passing 100-µl aliquots of cell-free extract from 200-ml cultures of overexpressed GCaMP2 over a Superdex 200 10/300 GL column while monitoring the absorption at 280 and 495 nm.

Spectrophotometric Analysis—Absorbance spectra were obtained in a Safire² (Tecan) with UVStar 96-well plates (Greiner) for both the calcium-free (10 mm EGTA) and calcium-loaded state (10 mm CaCl₂). For fluorescence spectra, Fluotrac 200 plates (Greiner) were used. Samples were diluted 10-fold in zero free calcium buffer (Invitrogen) (30 mm MOPS, 100 mm KCl, 10 mm EGTA, pH 7.2) for calcium-free spectra, and in 39 μM free calcium buffer (Invitrogen) (30 mM MOPS, 10 mм Ca-EGTA in 100 mм KCl, pH 7.2) for calcium-loaded spectra. For absorbance measurements, samples were dialyzed into 20 mm Tris, 100 mm NaCl, and EGTA or CaCl₂ was added to a final concentration of 10 mm.

RESULTS

Structural Analysis of Dimeric Ca²⁺-GCaMP2—To understand the molecular mechanism of Ca²⁺-dependent fluorescence signal change and facilitate efforts to improve GCaMP properties, we set out to determine structures of the calciumfree and calcium-bound states of GCaMP2. Initial efforts to crystallize affinity purified, calcium-saturated GCaMP2 led to a single crystallization hit, as described previously (17). The structure was solved by molecular replacement, and the final model revealed a domain-swapped dimer of GCaMP2 in the crystals, with the M13 peptide of each monomer bound by the

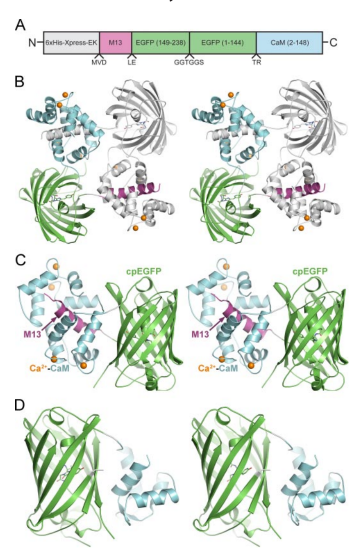


FIGURE 1. Crystal structures of GCaMP2. A, schematic of the primary amino acid sequence of GCaMP2 illustrating the domain organization. Domains are colored as depicted in B-D. Carets below the schematic show the positions of inter-domain linkers whose amino acid sequences are given. B, stereoview of the structure of the Ca²⁺-saturated domain-swapped GCaMP2 dimer, depicted as ribbons. One molecule of the dimer is colored by domain as in A, the other molecule is colored light gray. The EGFP chromophore is represented as sticks and calcium ions are shown as orange spheres. C. structure of Ca^{2+} -saturated GCaMP2 monomer, represented as in B except the domains are labeled. D, structure of calcium-free GCaMP2, represented as in B and C. Note that the M13 peptide and the C-terminal half of CaM are not included in the model due to lack of electron density, suggesting their flexibility. This and other structure figures were prepared using PyMOL (Delano Scientific, San

calcium-loaded CaM of the other (Fig. 1B and Table 1). The finding of a domain-swapped dimer structure was unexpected, given that GCaMP has previously been characterized as a monomer (9). We therefore characterized the oligomeric distribution of GCaMP2 in solution using several biophysical techniques.

Solution Characterization of GCaMP2-SEC analysis of affinity purified GCaMP2 in the absence of calcium showed a single peak close to the elution volume predicted for the monomeric form of GCaMP2, when compared with the elution volume of standard proteins (Fig. 2A). However, in the presence of saturating calcium, two peaks eluted. The two peaks were

TABLE 1Data collection and refinement statistics

	GCaMP2 dimer (PDB 3EK7)	GCaMP2 monomer (PDB 3EK4)	GCaMP2-T116V- K378W (PDB 3EKH)	GCaMP2-T116V- G87R (PDB 3EK8)	8EF-GCaMP2 (PDB 3EKJ)
Data collection					
Radiation source	APS 31-ID ^a	ALS BL8.2.2 ^b	APS 31-ID ^a	Copper anode ^c	APS 31 -ID ^a
Wavelength (Å)	0.9793	1.000	0.9793	1.5418	0.9793
Space group	C2	$P2_{1}2_{1}2$	$P4_{1}2_{1}2$	$P4_{1}2_{1}2$	C2
Cell dimensions					
a, b, c (Å)	126.13, 47.30, 68.94	60.49, 68.80, 117.26	121.64, 121.64, 97.32	120.82, 120.82, 97.35	211.87, 47.67, 42.99
α , β , γ (°)	90, 100.48, 90	90, 90, 90	90, 90, 90	90, 90, 90	90, 97.61, 90
Resolution (Å)	67.79-1.80 (1.90-1.80)	50.00-2.65 (2.74-2.65)	25.90-2.00 (2.11-2.00)	29.30-2.80 (2.90-2.80)	31.94-2.80 (2.95-2.80
$R_{\rm sym}$	7.6 (42.2)	7.5 (27.9)	9.6 (60.1)	18.4 (57.3)	15.4 (61.7)
$I/\sigma I$	18.8 (5.1)	20.44 (4.2)	21.5 (4.7)	8.6 (3.5)	15.4 (3.2)
Completeness (%)	98.5 (97.6)	98.7 (92.1)	100.0 (100.0)	100.0 (100.0)	98.9 (98.6)
Redundancy	7.5 (7.6)	3.1 (3.0)	14.1 (13.5)	11.94 (12.04)	7.1 (7.1)
Refinement					
Resolution (Å)	1.85	2.65	2.00	2.80	2.80
Unique reflections	33,937	14,650	49,718	18,310	10,580
$R_{\rm work}/R_{\rm free}$	0.189/0.241	0.222/0.280	0.194/0.224	0.224/0.266	0.210/0.283
No. atoms (AU)	3,333	2,799	3,472	3,182	2,430
Protein	3,097	2,767	3,185	3,154	2,425
Ligand/ion	4	3	10	4	
Water	198	29	277	24	5
B-factors $(\mathring{A}^2)^d$					
Protein	25.4	44.0	33.4	38.8	31.1
Ligand/ion	19.4	75.3	38.7	41.4	
Water	32.2	29.4	37.1	23.3	12.9
Root mean square deviations					
Bond lengtĥs (Å)	0.018	0.010	0.011	0.011	0.016
Bond angles (°)	1.735	1.336	1.332	1.335	1.691

^a APS 31-ID, Advanced Photon Source, Beamline 31-ID.

assigned as a monomeric form with a higher elution volume than the calcium-free monomer, and a smaller population of dimeric GCaMP2.

Analytical ultracentrifugation of GCaMP2 in the presence and absence of calcium gave results similar to SEC (Fig. 2B), as did dynamic light scattering (data not shown). These results indicate that calcium-bound GCaMP2 is more compact than calcium-free, consistent with data for calmodulin and M13pep (24, 28, 29). The observed dimeric structure, which preferentially crystallized in the initial screen, presumably represents the larger species in solution observed by SEC and analytical ultracentrifugation. After separation of the monomeric and dimeric calcium-saturated forms of GCaMP2 by SEC, neither showed significant presence of the other oligomeric state when re-analyzed days later, indicating that the kinetics of the monomer \leftrightarrow dimer equilibrium are very slow in the presence of saturating calcium.

The monomeric and dimeric forms of Ca²⁺-bound GCaMP2 have different absorption and fluorescence properties (Fig. 2*C*), suggesting an altered chromophore environment between the states. Monitoring absorbance at both 280 and 495 nm, as well as fluorescence at 488/510 nm during SEC separation of purified GCaMP2 monomer and dimer clearly indicated that the dimeric species had significantly lower chromophore absorption and fluorescence.

Fluorescence absorbance/excitation and emission scans of freshly separated monomeric and dimeric Ca²⁺-GCaMP2 indicated that apo, monomer, and dimer had nearly identical peak wavelengths for excitation and emission, but that the dimeric fraction was less fluorescent than the monomeric fraction, although both are substantially more fluorescent than the apo

fraction (Fig. 2*D*). Comparison of the absorbance spectra of the three states shows that the chromophore environment changes dramatically upon Ca²⁺ binding: the protonated non-fluorescent state of the GCaMP2 chromophore is diminished upon Ca²⁺ binding, indicated by the smaller band at 400 nm in both Ca²⁺-saturated states, explaining the mechanism of the calcium-dependent fluorescence increase. The presence of a larger proportional amount of protonated non-fluorescent chromophore in the dimeric fraction compared with the monomeric fraction is in agreement with the SEC data (Fig. 2*C*), indicating that the monomeric GCaMP2 species is brighter than dimeric GCaMP2.

IGCaMP2] in Neurons—Calcium imaging experiments may be adversely affected by contribution from two competing species in situ. We thus attempted to quantitate the monomerdimer equilibrium present at physiologically relevant GCaMP concentrations. Two-photon laser scanning microscopy of in utero electroporated GCaMP2 under control of the CAG promoter was used to estimate sensor concentration in an acute slice of mouse somatosensory cortex, by comparison with a concentration series of purified Ca²⁺-free GCaMP2 protein in thin-window cuvettes (supplemental Fig. S2). Under these conditions, intracellular [GCaMP2] was found to be ~10 μm. A previous report expressing GCaMP2 in transgenic mouse heart from a modified α myosin heavy chain promoter estimated the intracellular concentration to be 1.6 μm (13).

To understand what fraction of calcium-saturated GCaMP2 should be dimeric at these intracellular concentrations, we removed calcium from purified GCaMP2 protein with excess EGTA, prepared a dilution series of GCaMP2, added back excess calcium, and observed the proportion of monomeric and



^b ALS BL8.2.2, Advanced Light Source Beam Line 8.2.2.

^c Data collected on home source using Rigaku Rotating Copper Anode RUH3R.

^d B-factors were calculated on the STAN server using MOLEMAN2 (xray.bmc.uu.se/cgi-bin/gerard/rama_server.pl).

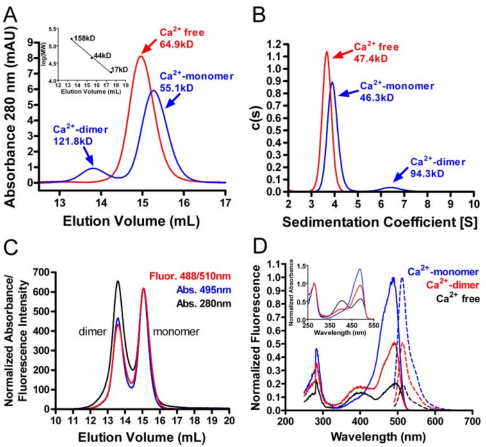


FIGURE 2. Solution studies of monomeric and dimeric GCaMP2. A, SEC of calcium-free (red) and calciumbound (blue) GCaMP2 following affinity purification. Peaks are labeled with the assigned species and estimated molecular mass. The standard curve used to estimate M_r is in the *inset*. The predicted molecular mass for GCaMP2 monomer and dimer are 50.7 and 101.3 kDa, respectively. B, c(s) distribution analysis of sedimentation velocity analytical ultracentrifugation data for calcium-free (red) and calcium-bound (blue) GCaMP2 following affinity purification. Peaks are labeled as in B. C, SEC analysis of a nearly equimolar mixture of purified, Ca²⁻ saturated GCaMP2 monomer and dimer, monitoring protein absorbance at 280 nm (black), cpEGFP chromophore absorbance at 495 nm (blue), and cpEGFP fluorescence at 488/510 nm (red). Note the difference in relative chromophore absorbance and fluorescence of the dimer. D, fluorescence excitation (solid lines) and emission (dashed lines) spectra of isolated Ca2+-saturated GCaMP2 monomer (blue) and dimer (red), as well as calcium-free GCaMP2 (black). Absorbance spectra of these three species are shown in the inset.

dimeric GCaMP2 using analytical SEC (supplemental Fig. S3). At the concentrations seen in the brain slice, only \sim 5% of the GCaMP2 was present as a dimer, suggesting that the monomeric sensor predominates under the conditions of typical imaging experiments.

Engineering Monomeric GCaMP2—Given that the Ca²⁺-GCaMP2 dimer is presumably not the predominant form under imaging experimental conditions, we attempted to determine the structure of the monomeric form of Ca²⁺-GCaMP2. Preparative separation of the Ca²⁺-GCaMP2 monomer by SEC and immediate crystallization screening led only to crystals of the dimeric form that grew after several weeks.

To aid crystallization of the monomeric form, we made a series of single amino acid substitutions in GCaMP2 to selectively disrupt dimerization, based on visual inspection of the Ca²⁺-GCaMP2 dimer structure. Eleven single amino acid substitutions in GCaMP2 were chosen and produced by site-directed mutagenesis (supplemental Table S1). Six of these mutations appeared to decrease the amount of GCaMP2 dimer present in purified protein samples as analyzed by SEC (supplemental Fig. S4). Sparse-matrix crystallization screening was carried out with these six GCaMP2 mutants, and two of them (GCaMP2 G87R (gfp G174R) and GCaMP2 K378W (cam K75W), where gfp refers to green fluorescent protein and cam refers to calmodulin) crystallized with a unique morphology under conditions distinct from those used for dimer crystallization (a topology diagram and amino acid sequence displaying both GCaMP and EGFP/CaM numbering are provided in supplemental Figs. S5 and S6, respectively). Although the G87R and K378W mutations appear to decrease dimer formation, the $\Delta F/F_0$ for each was not significantly altered (supplemental Table S1) suggesting that the mutations did not disrupt sensor function.

X-ray diffraction data were collected from these new mutant crystals, and the structure was determined by molecular replacement using the cpEGFP and CaM-M13 domains from the dimer structure. The relative orientations of the cpEGFP and CaM-M13 domains revealed that the quaternary organization was different from the dimer structure, and electron density corresponding to the inter-domain linkers clearly showed that the structure represented the monomeric form of Ca²⁺-GCaMP2.

Structural Analysis of Monomeric Ca²⁺-GCaMP2—Subsequently, crystals of the native Ca²⁺-GCaMP2 monomer were grown by seeding drops of Ca²⁺-GCaMP2 protein with microcrystals of the GCaMP2 K378W monomer. Solution of the Ca²⁺-GCaMP2 monomer structure by molecular replacement revealed that although these crystals were grown by seeding with K378W microcrystals, they were distinct from the G87R, K378W, and GCaMP2 dimer crystals (see Table 1).

The Ca²⁺-GCaMP2 monomer structure is very similar to that of the K378W and G87R mutants (root mean square deviations values of 0.7 and 0.6 Å, respectively, for comparing all 353 common C_{α} atoms), but the C-terminal lobe of the CaM domain is largely disordered in the Ca²⁺-GCaMP2 monomer crystals and could not be modeled well. Therefore, further discussion of the Ca²⁺-GCaMP2 monomer structure is based on analysis of the native Ca2+-GCaMP2 monomer structure together with the K378W and G87R structures, in which the CaM C-terminal lobe is better defined.

The domain organizations of the dimeric and monomeric forms of Ca²⁺-GCaMP2 are presented in Fig. 1, B and C. In the



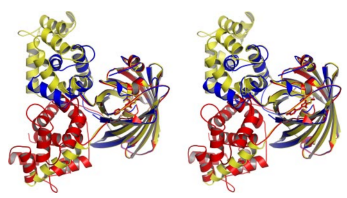


FIGURE 3. Stereoview of the structures of Ca²⁺-saturated GCaMP2-K387W monomer (red), Ca²⁺-saturated GCaMP2 dimer (yellow), and Ca²⁺-free 8EF-GCaMP2 (blue) superimposed using the cpEGFP domain. The proteins are represented as ribbons with the cpEGFP chromophore represented as sticks.

monomer structure, the Ca²⁺-CaM/M13pep complex makes more substantial contacts with the chromophore environment of cpEGFP, relative to the corresponding interaction in the dimer. Furthermore, the interaction surface of the Ca²⁺-CaM/M13pep with cpEGFP is quite different between the monomer and dimer (Figs. 3 and supplemental S7), with the former preferred in solution and the latter apparently favored by crystal packing.

As described (9), the EGFP domain of GCaMP is circularly permuted by connecting the native N and C termini with a flexible linker (GGTGGS), and by opening one of the β -strands of the EGFP barrel by elimination of amino acids 145-148 (supplemental Fig. S5), resulting in new N and C termini at the side of the barrel, directly abutting the EGFP chromophore. The GCaMP2 structures reveal that circular permutation of EGFP at this location has very little effect on the overall tertiary structure of the barrel or loops. Structural superimposition of the cpEGFP domain of either monomeric or dimeric GCaMP2 with the native GFP structures (PDB codes 1GFL (30) or 1EMA (31)) shows root mean square deviations of only \sim 0.4 Å when comparing all common 220 C_{α} atoms (supplemental Fig. S8A). The GGTGGS linker connecting the native EGFP N and C termini is disordered in all of the GCaMP2 crystals. The circular permutation of EGFP resulted, as expected, in an opening to the interior of the barrel due to the removal of four residues of the β -strand, although this opening is partially occluded in both Ca²⁺-bound structures by the Ca²⁺-CaM/M13pep complex. The opening measures \sim 5 Å \times 10 Å and allows appreciable solvent access to the *p*-hydroxybenzylideneimidazolinone chromophore. The phenolate moiety of the chromophore tyrosine points directly out of the opening, and is the most solventexposed portion of the chromophore. In the GCaMP2 dimer structure, large channels allow access from the surface of the protein, through this opening, to the chromophore. However, in the monomeric conformation the opening is mostly occluded by the "linker" connecting the N-terminal and C-terminal lobes of CaM, as well as the linker connecting cpEGFP with CaM. Solvent access is likely to lead to protonation of the phenolate oxygen (p $K_a \sim 6.0$ in EGFP and ~ 7.1 in GCaMP), which in EGFP is stabilized in the fluorescent, deprotonated form by a large hydrogen bond network. This may partially

explain the difference in calcium-saturated fluorescence between the monomer and dimer.

Each Ca²⁺-bound CaM-M13 complex present in both the monomer and the dimer Ca2+-GCaMP2 structures closely resembles the isolated CaM-M13 peptide complex structure determined previously (28), with the two lobes of CaM wrapping around and enveloping the helical M13 peptide. Structural superimposition of each CaM-M13 unit of GCaMP2 with this isolated human CaM-M13 complex structure (PDB code 1CDL) gives root mean square deviation values of 0.6 Å for the dimeric form, and 1.6 Å for the monomer when comparing all 161 common C_{α} atoms (supplemental Fig. S8B). The primary source of the larger structural deviation of monomeric GCaMP2 is the inter-lobe "linker helix" of CaM, which packs against the opening of the cpEGFP domain in the monomer structure but has fewer interactions in the dimer. This linker, which is known from previous structural work to adopt different conformations (24, 28, 29), is one α -helical turn longer in its N-terminal section and one turn shorter in the C-terminal section of the Ca²⁺-GCaMP2 monomer structure, relative to the dimer (supplemental Figs. S5 and S8B). This conformational change is a direct consequence of the interaction of the CaM domain with the cpEGFP domain, and has important implications for sensor functionality.

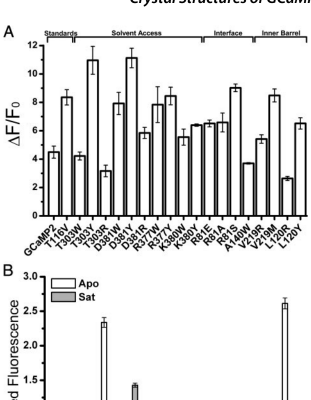
The cpEGFP domain is tethered in two ways to the calciumresponsive CaM in the calcium-saturated structures: the linker covalently connecting the C terminus of cpEGFP with the N terminus of CaM (cpEGFP-CaM linker), and the non-covalent interaction of CaM with M13pep, which is connected to the N terminus of cpEGFP by a short linker (M13-cpEGFP linker). Of these two linkers, the cpEGFP-CaM linker appears to be the more flexible, with ~5 consecutive amino acids between secondary structure elements or anchoring interactions, and with only weak electron density. The M13-cpEGFP linker, conversely, is shorter, less flexible, and makes several interactions with both the cpEGFP and CaM domains. In addition to the linkers and the CaM-M13 interaction, inter-domain contacts are made directly between cpEGFP and CaM. In the GCaMP2 monomer structure, the N-terminal lobe and inter-lobe linker of CaM contact the side of the cpEGFP barrel on opposite sides of the opening, occluding \sim 638 Å² of the solvent-accessible surface of each domain at the interface (supplemental Fig. S7). Very few interactions are made between cpEGFP and the C-terminal lobe of CaM in the monomer. In the dimer, each CaM domain contacts both cpEGFP domains (Figs. 1B and supplemental S7); the most extensive contact interface is observed between the N-terminal lobe of CaM and the side of the cpEGFP barrel of the same GCaMP monomer. Approximately 625 Å² of the solvent-accessible surface of each domain is buried at this interface. Two smaller patches of contact surface, totaling \sim 249 Å², are present between each lobe of the CaM domain and the side of the cpEGFP barrel from the other molecule of the GCaMP dimer.

Engineering and Structural Analysis of Calcium-free GCaMP2—The GCaMP sensor was designed on the assumption that in the absence of calcium, CaM would not associate with M13pep, allowing significant solvent access-mediated darkening of the EGFP chromophore (9). Understanding the mechanism of calcium-free fluorescence decrease could



allow targeted mutagenesis to improve baseline fluorescence or total dynamic range. To observe the domain organization of GCaMP2 in the absence of calcium (apo-GCaMP2) we attempted crystallization in the presence of an excess calcium chelator (EGTA). All crystallization trials produced dimeric, Ca²⁺-bound GCaMP2, despite repeated attempts to remove calcium from buffers, protein, glass, and plasticware. We therefore engineered an 8-fold mutant of GCaMP2, with two mutations in each of the four Ca2+-binding EF hands in the CaM domain (T329G/E334Q, D359G/E370Q, D396G/E407Q, D432G/E443Q) that should disrupt calcium binding. This mutant, 8EF-GCaMP2, spectrally resembles apo-GCaMP2 and does not respond to Ca^{2+} ($\Delta F/F_0 = 0.15 \pm 0.04$) (see supplemental Fig. S9 for spectra). Sparse-matrix crystallization screens with 8EF-GCaMP2 produced thin plate crystals that diffracted to 2.8 Å. The structure of 8EF-GCaMP2 was solved by molecular replacement using the cpEGFP domain from the previous structures, manual placement of the N-terminal lobe of apo-CaM into the remaining electron density, and rebuilding of the inter-domain linkers (Fig. 1D). No electron density was present for either the C-terminal lobe of CaM or the M13 peptide, indicating that in the absence of calcium the inter-domain flexibility is increased. The N-terminal lobe of CaM in 8EF-GCaMP2 closely resembles the "closed" calcium-free conformation observed in previous solution structures of apo-CaM (24, 29), with substantial changes in the inter-helical angles serving to sequester the hydrophobic target-binding cleft (Figs. 3 and supplemental S8). The first and last helices of the N-terminal lobe of CaM pack against the barrel of cpEGFP, creating nearly the same interface observed in the dimeric Ca²⁺-bound structure. In the absence of inter-domain interactions blocking the opening of cpEGFP, as observed in the calcium-saturated structures, solvent is allowed direct access to the EGFP chromophore, which appears to be stabilized in the protonated, non-fluorescent form (supplemental Fig. S9). The position of the M13 peptide is not constrained by tight binding to CaM; consequently no electron density is observed for the M13pep. However, the linker connecting M13pep with cpEGFP (Leu⁶⁰– Glu⁶¹) is observed and adopts a significantly different conformation in apo-GCaMP2 compared with the monomer and dimer GCaMP2 structures (supplemental Fig. S10). In the calcium-free structure, this linker is incorporated into the β-strand leading into cpEGFP; the side chain of Glu⁶¹ hydrogen bonds to the backbone amide nitrogen of Arg⁸¹, stabilizing this conformation (supplemental Fig. S10).

Structure-based Mutagenic Analysis of GCaMP2 Function— With the GCaMP2 crystal structures in hand, we set out to analyze GCaMP sensor function by site-directed mutagenesis to understand the mechanistic basis for the response to calcium. As GCaMP2 is largely monomeric under typical imaging conditions, the Ca²⁺-monomer structures served as the basis for mutagenesis studies to address function. All mutations were made in the context of GCaMP2-T116V (gfp T203V), a mutation previously described to increase the wild type EGFP excited-state proton transfer (32-34).3 We created groups of GCaMP2 mutants to



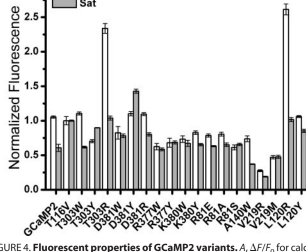


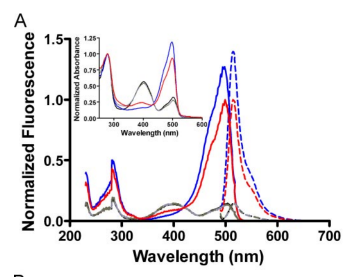
FIGURE 4. Fluorescent properties of GCaMP2 variants. A, $\Delta F/F_0$ for calcium binding of all GCaMP2 mutants listed in supplemental Table S3. All GCaMP2 mutants listed also contain the T116V mutation. B, normalized fluorescence of Ca²⁺-free (apo, white) and the Ca²⁺-saturated (Sat, gray) state of the GCaMP2 variants. Fluorescence of both states is normalized to the brightness of GCaMP2 T116V. Note that although GCaMP2 T116V-T303Y has a greater $\Delta F/F_0$ (panel A), the brightness of both Ca²⁺-free and Ca²⁺-saturated state is lower than for GCaMP2 T116V.

test three hypotheses about GCaMP function: (a) that mutating CaM or the cpEGFP-CaM linker to block solvent access to the EGFP chromophore would improve brightness ("solvent access" mutants), (b) disrupting the cpEGFP-CaM interfaces seen in the crystal structure would decrease sensor performance ("interface" mutants), and (c) mutating EGFP β -barrel positions observed to be solvent-exposed in the GCaMP2 structure (but not in EGFP itself) due to the circular permutation would alter sensor function ("inner barrel" mutants). Eighteen mutations, through the three mutant classes, were created by site-directed mutagenesis and analyzed by measuring fluorescence of purified protein (Figs. 4 and supplemental S9).

Two of the mutations predicted to block solvent access to the chromophore by introduction of a bulkier amino acid side chain near the cpEGFP opening were successful. GCaMP2-T303Y (in egfp-cam linker) and GCaMP2-D381Y (cam D78Y) showed significantly improved $\Delta F/F_0$ for Ca²⁺ binding (Fig. 4A). The D381Y (cam D78Y) mutation had a slightly brighter



³ L. Tian, S. Hires, J. Akerboom, E. Schreiter, and L. Looger, manuscript in preparation.



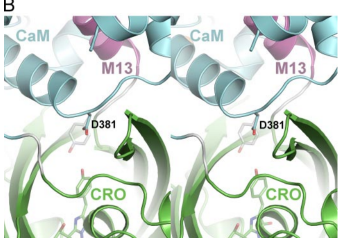


FIGURE 5. **A rationally designed, improved GCaMP2 variant.** *A*, fluorescence excitation (*solid lines*) and emission (*dashed lines*) spectra of Ca²⁺-saturated GCaMP2 T116V (*red*) and T116V/D381Y (*blue*), as well as their calciumfree forms (*gray* and *black*, respectively). Normalized absorbance spectra of each are shown in the *inset. B*, close-up stereo view of the Ca²⁺-saturated monomeric GCaMP2 structure, showing the location of aspartate 381 (*D381*) of CaM at the CaM/cpEGFP domain interface. GCaMP2 is displayed as *ribbons* colored by domain. The side chain of Asp³⁸¹ and the cpEGFP chromophore are displayed as *sticks*. A model of a tyrosine side chain at position 381 is shown in *semitransparent sticks* to represent a possible conformation of the D381Y mutant and to illustrate the proximity of this side chain to the chromophore.

apo state (Fig. 4*B*), a substantially brighter Ca²⁺-bound state, and showed the highest dynamic range of all mutants tested ($\Delta F/F_0$, D381Y = 11.1; $\Delta F/F_0$, GCaMP2 = 4.5; $\Delta F/F_0$, T116V = 8.4). Absorbance and fluorescence scans showed that D381Y (cam D78Y) has a lower proportion of protonated chromophore in the Ca²⁺-bound state (Fig. 5*A*), thus improving signal-to-noise ratio. Position Asp³⁸¹ of GCaMP2 is located on the flexible region of CaM between the two calcium-binding lobes and packs directly against the cpEGFP opening in the Ca²⁺-monomer structures (Fig. 5*B*). The side chain of a tyrosine at position 381 could extend into the opening of the cpEGFP barrel and come within a few angstroms of the chromophore (Fig. 5*B*).

Disruption of the cpEGFP-CaM interfaces observed in the Ca²⁺-GCaMP2 crystal structures dramatically decreased saturated-state fluorescence (Fig. 4). Position Arg⁸¹ (gfp Arg¹⁶⁸),

from a β -strand lining the cpEGFP opening, was observed to form salt bridges with Glu⁶¹ of the M13-cpEGFP linker and Glu³⁸⁷ (cam Glu⁸⁴) from the CaM inter-lobe linker, in addition to a hydrogen bond with the backbone carbonyl oxygen of Thr³⁸² (cam T79). Mutation of Arg⁸¹ (gfp Arg¹⁶⁸) to glutamate, alanine, or serine decreased Ca²⁺-saturated fluorescence by ~25%. Mutation of Ala¹⁴⁰ (gfp Ala²²⁷), near the center of the Ca²⁺-CaM/cpEGFP interface, to the bulky amino acid tryptophan resulted in a 60% decrease in Ca²⁺-saturated fluorescence. These disruptive mutations suggest that the domain interfaces observed in the Ca²⁺-monomer crystal structure are functionally relevant in solution as well.

Two positions in EGFP (gfp 62, gfp 207), normally tightly packed inside the barrel but partially exposed by circular permutation, were also selected for mutagenesis. Because both positions are near the chromophore, we hypothesized that mutations to larger side chains could significantly alter the chromophore environment. Mutation of Val²¹⁹ (gfp Val⁶²) to arginine or methionine dramatically decreased both apo and Ca²⁺-saturated fluorescence (Fig. 4B). Because Val²¹⁹ (gfp Val⁶²) is located only one turn of α -helix from the chromophore, it is possible that these mutations interfere with chromophore maturation or destabilize the cpEGFP structure. Mutation of Leu¹²⁰ (gfp Leu²⁰⁷) to arginine dramatically increased apo fluorescence without significantly altering Ca²⁺saturated fluorescence, whereas mutation to tyrosine increased apo fluorescence and slightly decreased Ca²⁺-saturated fluorescence. The mechanism by which L120R (gfp L207R) leads to a large increase in apo fluorescence is unknown, but the large functional variance of the two mutations tested suggest that this could be an interesting position for future mutagenic libraries.

DISCUSSION

The structures described here illustrate the molecular mechanisms by which the steric and electrostatic environment of the GCaMP2 chromophore is altered upon calcium binding. Large scale conformational changes that alter relative domain orientations serve to change the stereochemical functionality presented to the chromophore at the cpEGFP barrel opening, and to control the access of solvent molecules (consistent with Ca²⁺dependent p K_a decrease of the GCaMP chromophore (9)). The biophysical characterization in solution demonstrated that GCaMP2 is predominantly monomeric at physiological concentrations during imaging experiments in tissue cells, although the issue of dimerization remains relevant because future GCaMP mutations could inadvertently favor the dimeric form and complicate interpretation of imaging experiments. Earlier analysis of GCaMP (9) used protein concentrations too low and gel filtration of inadequate resolution to observe dimerization.

The crystal structures reveal some of the mechanisms by which mutations previously made to the GCaMP scaffold, both in the initial engineering and in later rounds of optimization, function to improve the sensor. The crystal structures of GCaMP2 demonstrate the close proximity of the M13pepcpEGFP linker to the EGFP chromophore relative to the cpEGFP-CaM linker, which is consistent with the reported



effect of mutation of these two regions on sensor response (9). During the original construction of GCaMP it was noted that positively charged side chains at position 61 within the M13cpEGFP linker abolished responsiveness to calcium, whereas serine or threonine on that position led to photoisomerization (9). In the calcium-free 8EF-GCaMP2 structure, Glu⁶¹ occupies the same location as His^{148} in EGFP. The protonated N δ atom of His¹⁴⁸ is critical for the hydrogen bond network stabilizing the deprotonated state of the chromophore of EGFP (35, 36); Glu⁶¹ in the calcium-free 8EF-GCaMP2 could fulfill the same role and might preferentially stabilize the protonated state contributing to the decreased fluorescence observed in calciumfree GCaMP2. In the calcium-bound structures, Glu⁶¹ is pulled away from the cpEGFP barrel opening due to the interaction of M13pep with CaM and forms a salt bridge interaction with Arg⁸¹ from the outside of the cpEGFP barrel.

Small-probe contact dot analysis (37) of the EGFP crystal structure (PDB 1EMA) shows that Val¹⁶³ is over-packed in the hydrophobic core, potentially clashing with Ile¹⁵². Mutation of EGFP-Val¹⁶³ to Ala in GCaMP1.6 resulted in a more thermostable sensor (15), and the crystal structures of GCaMP2 show that Ala⁷⁶ (gfp 163) eases this unfavorable interaction without creating a cavity. Mutation of EGFP-Ser¹⁷⁵ in GCaMP1.6 to glycine similarly improved thermostability; in addition to improving packing within the EGFP domain, this change may allow backbone relaxation to facilitate interaction with the CaM inter-domain linker, critical for shielding the chromophore from solvent quenching. The GCaMP1.6 mutation GFP-A206K, intended to minimize EGFP dimerization, has an unexpected effect on the GCaMP sensor, as this position actually interacts with the CaM domain, rather than the EGFP domain of another sensor molecule. In the dimeric state, GCaMP2-Lys¹¹⁹ (gfp 206) forms an electrostatic interaction with GCaMP2-Glu³⁰⁹ (cam 6) within the GCaMP2 monomer. In the monomeric state, a similar electrostatic interaction occurs with GCaMP2-Asp305 (cam 2). Thus the actual effect of this mutation in GCaMP1.6 may be to stabilize EGFP-CaM contacts within the GCaMP monomer, rather than to disfavor EGFP dimerization. The GCaMP2 mutation GFP-D180Y appears to improve the β -strand propensity of the position regardless of Ca²⁺ binding and EGFP-CaM association. All five structures show that GCaMP2-Tyr93 (gfp 180) is solvent-exposed, with few interactions within the EGFP domain and none with other sensor components. The mutation V251I (gfp 93) in GCaMP2 increases packing interactions between two adjacent β-strands, particularly with GCaMP2-Ile¹⁰¹ (gfp 188). No interdomain contacts are observed at position 251 in the five crystal structures reported here, consistent with the calcium-independent stabilization observed with this mutation (13).

We have explored only a limited panel of mutations based on inspection of the crystal structures reported here. Future efforts to analyze additional point mutations and mutagenic libraries at positions shown to be of interest based on the crystal structures should lead to even better genetically encoded calcium indicators with further enhanced signal-to-noise capable of improved neural activity imaging. Additionally, we anticipate that the structural information gained from this work on

GCaMP2 will aid rational design efforts to produce improved fluorescent protein-based sensors for other analytes.

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Note Added in Proof—During the review process of this manuscript, Wang et al. (Wang, Q., Shui, B., Kotlikoff, M. I., and Sondermann, H. (2008) Structure (Lond.) 16, 1817–1827) reported on the structures of monomeric and dimeric GCaMP2. The results of that study are consistent with those described here.

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