Characterization of Two Human Skeletal Calsequestrin Mutants Implicated in Malignant Hyperthermia and Vacuolar Aggregates Myopathy

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Background: Hereditary mutations (D244G and M87T) of skeletal calsequestrin have been associated with skeletal myopathies.

Results: The D244G mutation loses Ca$^{2+}$, resulting in structural instability. M87T inhibits calsequestrin's polymerization by altering the Casq1 dimer interface.

Conclusion: D244G is largely dysfunctional, whereas the Ca$^{2+}$-binding capacity of M87T is mildly reduced.

Significance: Altered characteristics of Casq1 mutants are congruent with their associated disease phenotypes.

ABSTRACT

Calsequestrin 1 (hCasq1) is the principal Ca$^{2+}$ storage protein of the sarcoplasmic reticulum of skeletal muscle. Its inheritable D244G mutation causes a myopathy with vacuolar aggregates, while its M87T “variant” is weakly associated with malignant hyperthermia (MH). We characterized the consequences of these mutations with studies of the human proteins in vitro. Equilibrium dialysis and turbidity measurements showed that D244G and, to a lesser extent, M87T partially lose Ca$^{2+}$ binding exhibited by wild type hCasq1 at high Ca$^{2+}$ concentrations. D244G aggregates abruptly and abnormally, a property that fully explains the protein inclusions that characterize its phenotype. D244G crystallized in low Ca$^{2+}$ concentrations lacks two Ca$^{2+}$ ions normally present in wild type, which weakens the hydrophobic core of Domain II. D244G crystallized in high Ca$^{2+}$ concentrations regains its missing ions and Domain II order, but shows a novel dimeric interaction. The M87T mutation causes a major shift of the α-helix bearing the mutated residue, significantly weakening the back-to-back interface essential for tetramerization. D244G exhibited the more severe structural and biophysical property changes, which matches the different pathophysiological impacts of these mutations.

To activate contraction of skeletal muscle fibers, up to 200 μmoles/L of Ca$^{2+}$ are released into the cytosol after an action potential. Ca$^{2+}$ is released from the sarcoplasmic reticulum (SR), an organelle of small volume where the ion is mostly bound to acidic proteins (reviewed in (1)). Calsequestrin (2) (protein and genes denoted here as Casq) is the most abundant and capacious Ca$^{2+}$-binding protein within the SR of both skeletal and cardiac muscle, where tissue-specific isoforms, Casq1 and Casq2, are respectively expressed. In skeletal muscle, ions bound to Casq1 constitute approximately 75% of the Ca$^{2+}$ released for activation of contraction (3).

Ca$^{2+}$ titrations of Casq in vitro feature multiple stages associated with progressive Casq polymerization (4,5). We have interpreted this observation, together with crystallographic studies, as evidence that Casq binds Ca$^{2+}$ cooperatively, which explains both its greater oligomerization
and greater Ca\textsuperscript{2+} binding capacity as Ca\textsuperscript{2+} concentrations increase. In vivo, inside the SR, Casq exists in linear ramified polymers, forming a dense network that fills the SR terminal cisternae (6,7). The structure of this network is entirely consistent with the lattice interactions observed in vitro (8). Polymerized in this way, Casq in living muscle appears to be in a state consistent with maximum storage capacity.

Other structural details suggest roles for Casq additional to Ca\textsuperscript{2+} storage (reviewed in (9)). The polymeric network inside the SR ends with slender pillars or tendrils that lead to the junctional membrane, near the mouth of the RyR channels (7). Casq1 binds between 40 and 60 moles of Ca\textsuperscript{2+} per mol, but in past crystallographic studies only up to 17 Ca\textsuperscript{2+} ions were identified, the majority of which appeared to be loosely bound, and consequently, diffusible (8). These features suggest that Casq polymers facilitate Ca\textsuperscript{2+} diffusion towards the open Ca\textsuperscript{2+} release channels, both by increasing the local concentration of diffusible Ca\textsuperscript{2+} ions and by adding one-dimensional directionality down the polymer, like a “calcium wire” (10,11). This mechanism, a form of diffusion enhancement by reduction of dimensionality (12), remains hypothetical.

Finally, Casq appears to have a gating role, specifically operating in the termination of Ca\textsuperscript{2+} release. This termination, which is essential for rapid contractile relaxation, requires fast closing of SR Ca\textsuperscript{2+} release (RyR) channels. Some evidence indicates that Casq is required for adequate channel closure in skeletal muscle (13,14), but the issue remains controversial (15).

Much of the interest in the properties of this protein stems from observations of linkage between its mutations and human disease (special cases of “couplonopathies”, or diseases of the couplon (19)). This association is especially clear in the heart, where at least 15 Casq2 mutations have been linked with a disease known as catecholaminergic polymorphic ventricular tachycardia (CPVT2, reviewed in (16)).

As for Casq1, its association with muscle disease rests on three observations: i) A syndrome similar to malignant hyperthermia (MH) has been reported for Casq1-null mice (17,18). The causation of MH by Casq1 absence is consistent with the observations in cardiac muscle, as both are mechanistically similar diseases of enhancement and loss of control of Ca\textsuperscript{2+} release (19), and CPVT2 is, in most cases, associated with severe deficits in the amount of protein present (16). ii) A missense mutation, D244G, is linked to a myopathy characterized by the presence of vacuoles containing SR protein inclusions (20). The patients experience muscle weakness and their cells show altered Ca\textsuperscript{2+} dynamics. iii) A second missense mutation, M87T, was found in 16 of 205 MH probands (21). M87T could not be formally linked to MH (it is clinically called a variant, rather than a mutation—the term we use here), but a mild association of mutation and disease was found (with odds ratio of ~2). Additionally, this mutation occurred in patients also carrying a MH-causative mutation of RyR1, at an unexpectedly high frequency. While the M87T mutation appears to cause mild malfunction per se, it might contribute to the disease phenotype when associated to RyR mutations. M87T also has relevance in the context of its polymerization-dependent Ca\textsuperscript{2+} storage functions, as replacing the highly conserved Met87 by Thr at the dimer interface should hamper dimerization (21).

Here, we report the first structural and biophysical characterization of D244G and M87T hCASQ1, compare them to wild type hCasq1 and propose mechanisms for how these mutations may contribute to the associated disease phenotypes.

METHODS

Site-Directed Mutagenesis—The D244G and M87T hCasq1 genes were generated by site-directed mutagenesis of the wild type hCasq1 gene (GenBank: AB277764.1) in pET30a and transformed into Rosetta(DE3)pLysS cells for protein expression.

Protein Expression and Purification—Rosetta(DE3)pLysS E. coli cells containing the Casq1 vectors were grown in LB media at 37 °C and induced by 0.5 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) to overexpress reaching an A\textsubscript{600} of 0.6. Following induction, the cells were harvested, suspended in 20 mM tris, 0.05 g/100 mL Na\textsubscript{3}p, pH 7.5, sonicated at 8000 rpm using a 450 Sonifier® (Branson Ultrasonics), until they reached apparent homogeneity. The resulting lysate was clarified by centrifugation at 20,000 × g and loaded onto a Toyopearl DEAE-650M (Tosoh Biosciences) column equilibrated with DEAE wash buffer (20 mM tris, 0.05 g/100 mL Na\textsubscript{3}p, pH 7.5, 500 mM Na\textsubscript{2}SO\textsubscript{4}). The resulting protein was eluted stepwise with 200 mM Na\textsubscript{2}SO\textsubscript{4} and 600 mM Na\textsubscript{2}SO\textsubscript{4}

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mL NaN₃, pH 7.5). hCasq1 eluted between a linear gradient of 12.5 to 25 % DEAE elution buffer (20 mM tris, 2 M NaCl, 0.05 g/100 mL NaN₃, pH 7.5). DEAE fractions containing hCasq1 were then purified using a CHT ceramic hydroxyapatite column (Bio-Rad Laboratories) equilibrated with HA wash buffer (5 mM sodium phosphate, 0.05 g/100 mL NaN₃, pH 6.8). hCasq1 eluted between a gradient of 50 to 100 % HA elution buffer (0.5 M sodium phosphate, 0.05 g/100 mL NaN₃, pH 6.8). Finally, HA fractions containing purified hCasq1 were purified using a phenyl sepharose 6 fast-flow high-sub column (GE Healthcare) equilibrated with phenyl sepharose wash buffer (20 mM MOPS, 0.5 M NaCl, 0.05 g/100 mL NaN₃, 1 mM EGTA). After extensive washing, hCasq1 was eluted from the column using wash buffer containing 10 mM CaCl₂. Fractions containing hCasq1 were then buffer-exchanged into Casq assay buffer (20 mM MOPS, 0.3 M KCl, 0.05 g/100 mL, pH 7.2). Protein concentrations were determined using the bicinchoninic acid (BCA) assay (Thermo Scientific).

**Crystallization**—Wild type, D244G, and M87T hCasq1 (12.5 mg/mL in 20 mM HEPES, 0.5 M NaCl, 0.05 g/100 mL NaN₃, pH 7.0) were crystallized using the hanging drop vapor diffusion method at 4 °C. For wild type and M87T, 1.5 µL of protein solution was mixed with 1.5 µL buffer of crystallization buffer (0.1 M HEPES, 0.2 M NaCl, 27.5 (v/v) % 2-methyl-2,4-pentanediol, pH 7.0), whereas for D244G, 1.25 µL of protein solution was mixed with 1.75 µL of crystallization buffer to obtain the low-Ca²⁺ form. Crystals generally formed within 1 week. In the case of M87T, crystals grew from protein incubated with 5 mM CaCl₂ prior to mixing with crystallization solution. High-Ca²⁺ D244G crystals were obtained by slowly diffusing 10 mM CaCl₂ across a semipermeable membrane to D244G over the course of 24 hours.

**Structure Determination**—Crystallographic data were collected at the Advanced Light Source Beamline 8.2.1 (ALS BL8.2.1) and reduced and scaled using HKL2000 (22). The low-Ca²⁺ wild type hCasq1 structure was solved by molecular replacement using PHENIX with native rabbit Casq1 (PDB: 3TRQ) as an input model. The D244G and M87T structures were then solved by molecular replacement using the low-Ca²⁺ wild type hCasq1 structure as the input model. The high Ca²⁺ forms were built manually, as necessary. Iterative model adjustment and refinement were completed using COOT (23) and PHENIX. Crystallographic coordinates and structure factors for wild-type hCasq1 (PDB ID: 5CRD), low-Ca²⁺ D244G hCasq1 (PDB ID: 5CRI), high-Ca²⁺ D244G hCasq1 (PDB ID: 5CRG), and M87T hCasq1 (PDB ID: 5CRH) have been deposited in the Protein Data Bank. Refinement statistics are listed in Table 2.

**Turbidity Assays**—The turbidity of hCasq1 solutions (i.e., absorbance at 350 nm) as a function of Ca²⁺ concentration was monitored using a Genesys 10S UV-Vis Spectrophotometer (Thermo Scientific). Assays were performed using 1.0 mL of 2.0 mg/mL wild type, D244G, and M87T hCasq1 in Casq assay buffer. Concentrated Ca²⁺ solutions (0.10 M, 0.25 M, 0.50 M, and 1.0 M) were added in 1.0 µL aliquots to the 1.0 mL Casq1 solutions in a quartz cuvette to achieve the proper calcium concentration. Upon addition of each concentrated Ca²⁺ aliquot, the samples were mixed by aspiration and allowed to equilibrate (dA350/dt = 0) before addition of the next aliquot. Dilutions from adding Ca²⁺ aliquots were included in data analysis. For descriptive purposes, the A350 as a function of Ca²⁺ concentration was least-squares fitted to the sum of two Boltzmann functions, namely:

\[
y = y_0 + \left( \frac{A_1}{1 + e^{-\frac{x-x_{01}}{k_1}}} \right) + \left( \frac{A_2}{1 + e^{-\frac{x-x_{02}}{k_2}}} \right) \quad \text{(Eq. 1)}
\]

Where y is A350; x is the Ca²⁺ concentration; y₀ is the initial value; A₁ and A₂ are the spans of the absorbance changes; x₀₁ and x₀₂ are the centers of the first and second transitions; and k₁ and k₂ are the first and second slope factors, respectively.

**Equilibrium Dialysis and Inductively-Coupled Plasma Optical Emission Spectrometry (ICP-OES)**—Equilibrium dialysis cells two adjoined 1.4 mL acrylic half-cells separated by a 4.0 cm² circular regenerated cellulose dialysis membrane (12-14 kDa cutoff). For each cell, 0.5 mL aliquots of either 10 µM wild type, D244G, or M87T hCasq1 in Casq assay buffer were added to one half-cell and solutions of varying CaCl₂ concentrations in Casq assay buffer were added to the other. The two solutions within each cell equilibrated for three days at room temperature on a rocking shaker. After equilibration, the 317.933
nm Ca\textsuperscript{2+} emission intensity of each sample was measured using a PerkinElmer Optima 3200 RL ICP-OES. The difference between the Ca\textsuperscript{2+} contained within the protein-containing and protein-free sides of a given cell determined the bound Ca\textsuperscript{2+} (i.e., Ca\textsuperscript{2+}{protein} - Ca\textsuperscript{2+}{no protein} = \Delta \text{Ca}^{2+}), which, when divided by its corresponding moles of protein, gave the fractional occupancy. The vertical bars at each point are standard deviations of three independent measurements. The fractional occupations versus their corresponding free Ca\textsuperscript{2+} concentrations were least-square fitted to the sum of two Hill equations:

\[ f = B_{\text{max},1} \left( \frac{\left[ \text{Ca}^{2+} \right]^{n_1}}{\left[ \text{Ca}^{2+} \right]^{n_1} + K_{d_1}} \right) + B_{\text{max},2} \left( \frac{\left[ \text{Ca}^{2+} \right]^{n_2}}{\left[ \text{Ca}^{2+} \right]^{n_2} + K_{d_2}} \right) \quad (\text{Eq. 2}) \]

Where \( f \) is fractional occupancy, \( B_{\text{max},j} \) (with \( j = 1 \) or 2) are the maxima of the two binding components, \( K_{d_j} \) are their corresponding dissociation constants, and \( n_j \) are their Hill coefficients.

**Quantum Mechanics: Molecular Mechanics Optimization and Electrostatic Potential Surface Generation**—The structures of wild-type human, low-Ca\textsuperscript{2+} D244G, and high-Ca\textsuperscript{2+} D244G Casq1 were prepared for QM:MM calculations using the PDB Prep Wizard in Schrödinger Maestro (24). The high-affinity site C Ca\textsuperscript{2+}-binding sites in each structure were then optimized in Gaussian 09 using the ONIOM (QM:MM) method with AMBER used for the low layer (25). The B3LYP level of theory with double-zeta correlation-consistent basis sets (cc-pVDZ for H and C; aug-cc-pVDZ for N and O; and cc-pwCVTZ for Ca\textsuperscript{2+}) was used for the high layer (26,27). Single-point calculations at the B3LYP level of theory with triple-zeta basis sets (cc-pVTZ for H and C; aug-cc-pVTZ for N and O; and cc-pwCVTZ for Ca) were then performed on each optimized structure. Self-consistent field total electron density and electrostatic potentials were generated from the single-point calculation at 12 and 6 points per bohr, respectively. Electrostatic potential surfaces were generated by plotting the electrostatic potential on its corresponding electron density at an isovalue of 0.0200 electrons/bohr\textsuperscript{3} in GaussView 3.09 (28).

**RESULTS**

We compared the Ca\textsuperscript{2+}-dependent polymerization properties of the wild type protein and two mutants by light scattering and turbidity measurements. We also compared their equilibrium Ca\textsuperscript{2+} binding properties, as well as their crystal structures, obtained with or without Ca\textsuperscript{2+} in the medium.

**Ca\textsuperscript{2+}-Dependent Polymerization**—The size-exclusion chromatography profiles of wild type, D244G, and M87T hCasq1 (Fig. 1) show that the three proteins are all monomers in 0 mM Ca\textsuperscript{2+} buffer (Fig. 1A) and have roughly the same response to rising Ca\textsuperscript{2+} concentrations at 1.0 and 1.5 mM Ca\textsuperscript{2+} (Fig. 1B and 1C). There appeared to be a small presence of wild type and D244G Casq1 tetramers at 1.5 mM Ca\textsuperscript{2+} (Fig. 1C inset); because of the low absorbance of each peak (implying a very small amount of protein in this state), the significance of this observation is not clear.

**Turbidity Assays**—Above 1.5 mM Ca\textsuperscript{2+}, monitoring Ca\textsuperscript{2+}-dependent polymerization of Casq1 by MALS was not possible because the higher order Casq1 polymers clogged the column matrix. To circumvent these limitations, Casq1 aggregation beyond 1.5 mM Ca\textsuperscript{2+} was monitored by turbidity assays. Wild type hCasq1 (solid line in Fig. 2) started a major transition at 1.4 mM Ca\textsuperscript{2+} and saturated by 9 mM Ca\textsuperscript{2+} with an absorbance of 0.2. D244G (dotted line) underwent a large increase in turbidity that started at 1.4 mM Ca\textsuperscript{2+} and ended at 4.4 mM Ca\textsuperscript{2+} with a final absorbance of 1.0. The increase in absorbance for M87T (dashed line) was substantially lower than wild type and D244G. Expecting to enhance the response, M87T was studied at 2.0 mg/mL and 4.0 mg/mL (filled and open squares, respectively, in Fig. 2). The 4.0 mg/mL solution attained a maximum turbidity 2.8 times higher than the 2.0 mg/mL maxima, but both still required high Ca\textsuperscript{2+} concentrations to become turbid, and neither became as turbid as wild type.

**Equilibrium Ca\textsuperscript{2+}-Binding Studies**—Equilibrium Ca\textsuperscript{2+}-binding curves calculated from equilibrium dialysis/ICP-OES data are shown in Fig. 3. The binding data were fit to the sum of two "Hill" functions (Eq. 2). The best-fit parameters are listed in Table 1.

Each curve features two distinct binding stages, with roughly sigmoidal Ca\textsuperscript{2+} concentration
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dependence. The mutants have lower Ca\(^{2+}\) binding capacity, and in both cases, the second binding stage occurs at a greater Ca\(^{2+}\) concentration than wild type. In all cases, the second binding stage has a higher \(K_d\) than the first stage and, most notably, a very high Hill coefficient, suggestive of high cooperativity in this stage. While each Casq1 features the two-stage binding property, the high-Ca\(^{2+}\) high-cooperativity component is most prominent in wild type, and decays in size (i.e., \(B_{\text{mu2}}\)) for both mutants, especially D244G. Both mutants also have a greater \(K_d\) and a much greater Hill coefficient of the second component (the one with highest cooperativity).

A final detail worth noting: At 0 mM Ca\(^{2+}\), both wild type and M87T had an average fractional occupancy of 4 mol Ca\(^{2+}\)/mol hCasq1, whereas D244G had only 2, corresponding to the number of Ca\(^{2+}\) observed per monomer in its low Ca\(^{2+}\) crystal structure (described below).

Crystal Structure of Wild Type hCasq1 —

Wild type hCasq1 crystallized in absence of added Ca\(^{2+}\) in the C222\(_1\) space group with one molecule in the asymmetric unit, yielding 2.08 Å resolution. Like the published rabbit Casq1 (PDB: 3TRQ) crystallized in the same condition, wild type hCasq1 contained Ca\(^{2+}\) ions at high affinity sites A, B and C, and a fourth Ca\(^{2+}\) bound by Thr189 (Fig. 4A). Wild type hCasq1 and rabbit Casq1 had similar structures, as demonstrated by the small root-mean-square deviation (r.m.s.d.) of their least-squares superposition (0.51 Å). In addition, the major lattice-packing interactions observed in both Casq1 crystals were mostly those involved in stabilizing a long linear polymer (Fig. 5A). The similarities of Casq1 between the two species, in terms of tertiary and polymeric structures, highlight Casq1’s conserved nature.

Crystal Structures of D244G hCasq1 —

Diffraction-quality crystals of D244G grew in both the absence of added Ca\(^{2+}\) and in the presence of 10 mM Ca\(^{2+}\), producing what we here call the low- and high-Ca\(^{2+}\) forms of D244G. Low-Ca\(^{2+}\) D244G crystallized in the P2\(_1\)2\(_1\)2\(_1\) space group with one molecule in the asymmetric unit, two Ca\(^{2+}\) ions per Casq1 monomer, and at a resolution of 3.32 Å (Fig. 4C). High-Ca\(^{2+}\) D244G crystallized in the P2\(_1\) space group with four molecules in the asymmetric unit, 21 Ca\(^{2+}\) ions per Casq1 monomer, and at a resolution of 1.97 Å (Fig. 4D).

Low Ca\(^{2+}\) D244G had a vacant and highly disordered high-affinity Ca\(^{2+}\)-binding site C (Fig. 6A right) presumably due to the mutation, which destabilized the hydrophobic core of Domain II (Pro160 through Thr263). Also affected were regions of Domain III connected to Domain II through high-affinity site B, a site responsible for attaching Domain II to III and located only a short stretch away from site C. The high-Ca\(^{2+}\) D244G structure, however, had bound Ca\(^{2+}\) at site C and, as expected, a well-ordered tertiary structure. Backbone atomic displacement parameter (ADP) plots confirmed these observations (Fig. 7). In the low-Ca\(^{2+}\) D244G structure, Domain I had similar order to wild type (i.e., similar ADPs), but had high disorder in Domain II, as well as in areas of Domain III connected to Domain II. In contrast, the high-Ca\(^{2+}\) D244G displayed a highly ordered structure throughout.

Each high-Ca\(^{2+}\) D244G dimer features intramolecular and intermolecular Ca\(^{2+}\) coordination at site C, with one monomer coordinating intramolecularly, and the second coordinating intermolecularly. The intramolecular coordination is similar to that of site C in wild-type hCASQ1, except for an additional solvent molecule (for a total of four) that is seen coordinating Ca\(^{2+}\) in the high-Ca\(^{2+}\) D244G structure. The intermolecular coordination at C, on the other hand, is unique to D244G. As shown in Fig. 8A, the Ca\(^{2+}\) bound by site C from a monomer in one dimer is coordinated by Glu360 from a monomer in a second dimer. This intermolecular coordination of site C forms non-canonical Casq1 tetramers (Fig. 8B). In each of the two dimers depicted, intramolecular coordination occurs on the Glu360-donating monomer. An intermolecular site C appears to be present in each dimer and causes this intermolecular coordination to propagate through the crystal lattice, making this interaction the possible cause for the Ca\(^{2+}\)-dependent crystalline aggregation of D244G seen in experiments.

To rationalize the different coordination modes, we analyzed the electrostatic potential surfaces of high-affinity site C from wild-type human and high-Ca\(^{2+}\) D244G. In wild type human Casq1 (Fig 8C), the carboxylate sidechain of Asp244, due to its proximity to the Ca\(^{2+}\) ion in site C, neutralizes the +2 charge conferred by the Ca\(^{2+}\) ion bound at site C. In the high-Ca\(^{2+}\) D244G
structure, site C coordinated intramolecularly (Fig. 8D) is left with a +1 charge as a consequence of losing the Asp244 carboxylate sidechain to the mutation. As shown by the electrostatic potential surface, this +1 charge is distributed throughout the Ca$^{2+}$-binding residues (here, Gly244, Pro246, Glu251, and four water molecules). In the case of the intermolecular coordination of site C by Glu360 from a monomer in a second dimer (Fig. 8E), direct Ca$^{2+}$ coordination by Glu360 neutralizes the +1 charge on site C. Two coordinating water molecules are displaced in the process, giving a final coordination sphere constituted by Gly244, Pro246, Glu251, Glu360, and two water molecules. As shown in Fig. 8, the intermolecularly-coordinated Ca$^{2+}$ ion is more neutralized than in wild-type, probably because the intermolecular case has two carboxylates directly coordinating Ca$^{2+}$ (Glu251 and Glu360), whereas wild-type has one direct coordination (Glu251) and one outer sphere anion (the Asp244 sidechain).

Crystal Structure of M87T hCasq1—M87T crystallized in the P2$_1$ space group with two molecules in the asymmetric unit. Crystals were obtained only in the presence of 2.5 mM Ca$^{2+}$ and each monomer contained 13 Ca$^{2+}$. The crystal structure had a resolution of 2.03 Å, which allowed clear imaging of the mutated residue (right side figure in Fig. 6B).

Least-squares superposition of the M87T structure onto the wild type hCasq1 structure, crystallized without adding Ca$^{2+}$, gave a r.m.s.d of 1.6 Å. This high deviation could have originated from the mutation, Ca$^{2+}$-dependent structural changes, or both. To separate these factors, we compared the bovine Casq1 crystal structure (PDB ID: 4TLY), obtained previously at 2.5 mM Ca$^{2+}$, with M87T hCasq1 (Fig. 9A). Their superposition had a lower r.m.s.d of 0.74 Å. The corresponding difference between the superposed structures, which was due to the mutation and not Ca$^{2+}$, was an outward shift of α-helix 2 (α2) in Domain I, where M87 is located (Fig. 9B).

DISCUSSION

We compared physicochemical properties of wild type hCasq1 and two naturally occurring mutants, D244G and M87T. D244G causes a myopathy associated with limited muscle function and characterized by vacuoles with inclusions consisting in aggregates of SR proteins. M87T has an ill-defined nosologic status. Its allele frequency in a large group of patients diagnosed as MH-susceptible (MHS) was greater than in control groups, but the difference was not significant (21). In spite of this inconclusive result, there are multiple indications that the mutation alters muscle function to some degree. Indeed, its presence increases the odds of MH by a factor of two. Furthermore, M87T was associated with causative RyR1 mutations in patients with the disease at a frequency higher than the expected random association value, which suggests that it enhances the disease phenotype.

Simultaneous Alteration of Ca$^{2+}$-Dependent Polymerization, Precipitation, and Ca$^{2+}$-Binding Capacity—At the free in vivo SR Ca$^{2+}$ concentration of -1 mM, CASQ1 is known to be fully polymerized, as shown by EM images of the junctional SR (6, 7). Although there are technical limitations to imitating an in vivo scenario, where CASQ1 concentrations approach 100 mg/mL and exists together with a number of other cellular components and structural constraints, our turbidity data confirmed that at least in vitro, D244G and M87T displayed different Ca$^{2+}$-dependent aggregation, in both quality and quantity, than that observed in the wild type. Although all forms showed similar monomer-to-dimer oligomerization between zero (Fig. 1A) and 1.0 mM Ca$^{2+}$ (Fig. 1B), wild type began to polymerize at lower Ca$^{2+}$ concentrations than either D244G or M87T (Fig. 2). However, D244G overtook wild type and M87T by 4 mM, reaching its maximum turbidity and presumably, maximum aggregation, by 4.4 mM Ca$^{2+}$ (Fig. 2). D244G precipitation was unique in that it produced by far the greatest increase in turbidity; furthermore, it settled rapidly at the bottom of the cuvette. The differences imply that the Ca$^{2+}$-induced quaternary change induced in D244G is qualitatively different, leading to larger aggregates. As argued below, these aggregates are less able to bind Ca$^{2+}$.

The M87T mutant responded less than the other Casq1 forms to rising Ca$^{2+}$ concentrations. The increase in turbidity only started around 5 mM Ca$^{2+}$ and reached a lower maximum, even at double the protein concentration (Fig. 2).

For all three Casq1s, the Ca$^{2+}$ binding data (Fig. 3) showed a binding process of roughly two
successive stages. The first stage corresponds largely to the occupancy of sites on individual protein monomers, a process of low cooperativity, which occurs at low Ca$^{2+}$ concentrations. Two aspects of the second binding stage, the requirement for higher Ca$^{2+}$ concentrations (reflected in a higher $K_d$) and the high Hill coefficient ($n_2$) are in agreement with the second stage consisting of Ca$^{2+}$ binding to polymerized Casq1, as multiple Ca$^{2+}$ ions are required to form the polymers. The second stage, at least in the wild type, has the greatest binding capacity, which explains the observed loss of Ca$^{2+}$ buffering power in vivo as the SR is depleted (1-3).

Both mutations resulted in a reduction of the total Ca$^{2+}$-binding capacity, mostly in the second stage (a reduction of $B_{\text{max2}}$), an increased $K_d$, and a large increase in $n_2$. These changes suggest that the highly cooperative second stage, which appears to be associated with Casq polymerization, is both more difficult to reach for the mutants and less effective as a means to provide additional binding capacity.

The mutation-dependent changes to the Ca$^{2+}$-binding properties of the two mutants were similar to the observed differences in the Ca$^{2+}$ concentration-dependent increase in turbidity. The Ca$^{2+}$-dependent increase in turbidity had the lowest steepness in M87T and required Ca$^{2+}$ concentrations much higher than wild type to reach its maximum turbidity. Both changes are consistent with the mutant’s reduced $B_{\text{max2}}$ and increased $K_d$. For D244G as well, the Ca$^{2+}$-dependent increase in turbidity shifted to higher Ca$^{2+}$ concentrations, in agreement with the change in $K_d$.

Overall, the quantitative and qualitative differences between D244G and wild type are probably causative for the vacuolar inclusion phenotype of vacuolar aggregates myopathy. Impaired targeting might be an additional cause of the functional deficits, as other important SR proteins were also detected in the vacuolar inclusions characteristic of this disease (21).

In agreement with the more overt disease phenotype linked to D244G, the physicochemical alterations appear greater in the D244G mutant than in the M87T variant. Other changes due to mutation, including the lower Ca$^{2+}$ binding capacity of both mutants, could explain some of the functional impairment observed in patients who have the diseases associated with these Casq1 mutations.

**Mutation-Dependent Changes to Protein Structure and Oligomeric Interactions**—In wild-type Casq1, including Casq1 from humans, the backbone carbonyl oxygen atoms of Asp244 and Pro246, together with the carboxylate of Glu251 and three water molecules, form a Ca$^{2+}$-coordination site known as high-affinity site C (Fig. 6A) (8). In the D244G mutant, the inherent flexibility of Gly244 destabilized site C and left it vacant, which caused an order-to-disorder transition not only for the immediate neighbors of the mutated residue but also for the adjacent hydrophobic core of Domain II. These disordered regions regained order when Ca$^{2+}$ concentrations rose sufficiently high enough for Ca$^{2+}$ to bind into the site C of D244G (Fig. 7). These observations confirm the previously hypothesized structural role of high-affinity site C.

The low-Ca$^{2+}$ D244G crystal lattice lacked the linear back-to-back interaction seen in wild type Casq1 (Fig. 5A), while the front-to-front interaction was relatively unchanged (Fig. 5B). Furthermore, the normal back-to-back interface did not reappear after D244G regained structural integrity under high Ca$^{2+}$ conditions (Fig. 5C). Instead, high-Ca$^{2+}$ D244G displayed a unique dimer-dimer interaction, mediated by intermolecular coordination of Ca$^{2+}$ at high-affinity site C. This intermolecular coordination would likely disrupt normal Ca$^{2+}$-dependent polymerization in vivo. The disordered and inadequately polymerized low-Ca$^{2+}$ D244G structure, taken together with the intermolecular coordination at site C in the high-Ca$^{2+}$ structure, shows that on a molecular level, the consequence of the D244G mutation is interference with the normal modes of action of high-affinity site C.

The Met87T mutation, at its core, is a disruption of the hydrophobic interactions that strengthen the front-to-front dimerization of Casq1. Met87 is located in Domain I on α-helix 2 (α2), which associates with α2 of its dimeric partner in complementary fashion within the front-to-front dimer interface (Fig. 9A). The hydrophobic residues on and near α2 form a hydrophobic pocket that plays a key role in dimerization by associating Met87 from the dimeric partner in a symmetric, reciprocal interaction. Because of the polar, hydrogen-
bonding nature of the substituted Thr residue, the M87T mutation inhibits closure of the hydrophobic pocket under low Ca\(^{2+}\) concentrations.

Upon exposure to rising Ca\(^{2+}\) concentrations, Domain I of Casq1 undergoes a large Ca\(^{2+}\)-dependent conformational change that exposes the hydrophobic face of a2 to solvent, with concomitant rotation of monomers around the front-to-front dimer interface. In M87T, the monomers rotated 3° less than wild-type bovine Casq1 under the same Ca\(^{2+}\) conditions (Fig. 9B). It is most likely that Thr87 can hydrogen bond with neighboring residues on a2 to form a more rigid structure, which may hamper Ca\(^{2+}\)-dependent rotation around the front-to-front interface. Supporting these expectations, instead of the linear lattice packing observed in the wild type (Fig. 5A), the crystal lattice of M87T was established mainly through side-to-side interactions between tetramers (Fig. 5D).

The altered front-to-front interface of M87T renders the dimer unsuitable for the linear back-to-back tetramerization that occurs with the wild type. If these non-linear tetramers were present in vivo, they would hinder further Ca\(^{2+}\)-dependent polymerization. A predictable consequence would be a reduced ability of the SR to store and release Ca\(^{2+}\) ions.

**Conclusion**—Overall, these missense mutations measurably alter the Ca\(^{2+}\)-dependent properties of Casq1. The D244G Casq1 mutant aggregates quickly and abnormally in response to rising Ca\(^{2+}\) concentrations, which explains the main pathological feature of the associated disease. The M87T Casq1 mutant, on the other hand, features lower reactivity to rising Ca\(^{2+}\) concentrations, probably due to an inability to polymerize to its full physiologically relevant extent. Since these properties—Ca\(^{2+}\) buffering and polymerization—are necessary for Casq1 to fulfill its role in the SR Ca\(^{2+}\) release/reuptake cycle, the D244G and M87T variants should lead to functional impairment. A reduction of the advantage for Ca\(^{2+}\) diffusion that calsequestrin “wires” putatively provide would also be expected. That they are most severe in the case of D244G, the most pathogenic mutation, indicates that its identified alterations are causative in the pathogenic process of vacuolar aggregates myopathy.

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**Author Contributions:** CK and ER conceived and coordinated this study. KML performed the biochemical/biophysical work, computational chemistry, and structure determination. LAR assisted with biochemical/biophysical work and structure determination. All authors contributed to data interpretation, the writing of this manuscript, and approved of all content contained herein.

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Abbreviations: Casq, calsequestrin; Casq1, skeletal calsequestrin; Casq2, cardiac calsequestrin; CPVT2, polymorphic ventricular tachycardia; D244G, Asp244-to-Gly244 mutation; ICP-OES, inductively-coupled plasma optical emission spectrometry; M87T, Met87-to-Thr87 mutation; MH, malignant hyperthermia; PDB, Protein Data Bank; RyR, ryanodine receptor (Calcium-release channel); SR, sarcoplasmic reticulum.
FIGURE LEGENDS

FIGURE 1. Multi-Angle Static Light Scattering. Multi-angle static light scattering profiles for wild type (solid line), D244G (dotted line), and M87T (dashed line) in (A) 0 mM Ca\(^{2+}\), (B) 1.0 mM Ca\(^{2+}\), and (C) 1.5 mM Ca\(^{2+}\). The inset in 1C is an expanded view that shows the low absorbance peaks at 1.5 mM Ca\(^{2+}\) between 5.5 and 7.5 mL, which correspond to the tetramer.

FIGURE 2. Ca\(^{2+}\)-Dependent Turbidity. Ca\(^{2+}\)-dependent turbidity curves for wild type hCasq1 (solid line, ●), D244G (dotted line, ♦), 2 mg/mL M87T (dashed line, ■), and 4 mg/mL M87T (dashed line, □). The inset is a magnified partial view of the same curves.

FIGURE 3. Equilibrium Dialysis Ca\(^{2+}\)-Binding Curve. Equilibrium dialysis-ICP-OES Ca\(^{2+}\)-binding curve, plotted as fractional occupancy (moles of bound Ca\(^{2+}\) per mole of hCasq1) versus free Ca\(^{2+}\) (mM) for wild type hCasq1 (solid line, ●), D244G (dotted line, ♦), and M87T hCasq1 (dashed line, ■). Bars represent standard deviations of three independent measurements.

FIGURE 4. hCASQ1 Crystal Structures. Monomeric structures of (A) low-Ca\(^{2+}\) wild type hCasq1, (B) high-Ca\(^{2+}\) M87T hCasq1, (C) low-Ca\(^{2+}\) D244G hCasq1, and (D) high-Ca\(^{2+}\) D244G hCasq1. Cyan and pink spheres represent Ca\(^{2+}\) bound at high-affinity and low-affinity sites, respectively. Note four Ca\(^{2+}\) in A, two in B, 21 in C and 13 in D. The small red circles and red triangles located on each structure indicate the N- and C-termini, respectively, to the furthest identifiable extent. The three high-affinity Ca\(^{2+}\) sites are boxed and domains I-III are circled in the wild-type structure (A).

FIGURE 5. Crystal Packing. Crystal packing and polymeric structure of (A) wild-type hCasq1, (B) low-Ca\(^{2+}\) D244G hCasq1, (C) high-Ca\(^{2+}\) D244G hCasq1, and (D) M87T hCasq1. Individual hCasq1 monomers are represented as their van der Waals surface and colored either blue or green. The asymmetric unit for each protein is circled.

FIGURE 6. Mutation Site Electron Densities. Crystallographic electron density maps of the D244G (A, right) and M87T mutation sites (B, right), corresponding to residues 240 through 259 and 84 through 98, respectively, compared to wild-type hCasq1 (A, left and B, left). Each electron density map (blue mesh) is a feature-enhanced map generated in PHENIX at a contour of 1.5 σ. The locations of mutated residues and their corresponding wild-type residues are indicated.

FIGURE 7. Atomic Displacement Parameter Plot. A plot of wild type (black solid line), low-Ca\(^{2+}\) D244G (black dashed line), and high-Ca\(^{2+}\) D244G (gray dashed line) main chain atomic displacement parameters (ADP) normalized to the mean wild-type chain value. The regions corresponding to Domain I – III and their limits at Pro160 and Thr263 are marked with vertical dashed lines. Residue 34 marks the N-terminus of Casq1 in its physiologically relevant form (i.e., after the signal peptide is removed).

FIGURE 8. Ca\(^{2+}\) Coordination in D244G. (A) Intermolecular Ca\(^{2+}\) coordination at high-affinity site C of D244G. The Ca\(^{2+}\) ion at site C is shown as a large cyan sphere and the small red spheres are coordinating water molecules. The green protein contains site C, and the gray protein contributes Glu360 for intermolecular coordination. Blue and red segments in each protein represent nitrogen and oxygen, respectively. The electron density map is contoured at 1.5 σ. (B) Dimers of D244G have two asymmetrical high-affinity site Cs: open circles mark intermolecularly coordinated site Cs, which support non-canonical tetramerization. Thus, site C in the green monomer (whose dimeric partner is in light purple) is complemented by Glu360 from the gray monomer (whose dimeric partner is in light blue). The dashed circles indicate high-affinity site Cs that are coordinated solely intramolecularly. Electrostatic potentials for wild-type hCasq1 site C (C), high-Ca\(^{2+}\) D244G intramolecular (D), and intermolecular site C (E). The potential surfaces are shown at an isovalue of 0.200 electrons/bohr\(^3\). The color scale
corresponds to a range of -0.140 kT/e (red) to +0.350 kT/e (blue), with green corresponding to a potential of 0.000 kT/e (or -0.200 for red to +0.500 in hartrees for blue, where 1 hartree = 627.509 kcal/mol).

**FIGURE 9. M87T Dimeric Interactions.** (A) M87T front-to-front dimer. The locations of α-helix 2 and Thr87 from each monomer are indicated. The gray helices represent the least-squares superposed α2 from high-Ca²⁺ wild type Casq1. (B) Ca²⁺-induced rotation of dimers around front-to-front interface of M87T (left) and wild type Casq1 (right). The solid and dashed lines represent the relative orientations of Domain I from each of the dimeric partners. The angles of rotation from the initial relative rotation of 180° across the front-to-front dimer interface are listed.
Table 1. Best-fit parameters for Ca\textsuperscript{2+} binding curves fit to a “Hill function”

|                | wild-type | D244G | M87T |
|----------------|-----------|-------|------|
| $K_{d1}$ (mM)$^a$ | 0.56      | 1.01  | 2.50 |
| $K_{d2}$ (mM)$^a$ | 3.60      | 4.35  | 4.70 |
| $B_{\text{max},1}$\textsuperscript{c} | 23.3      | 37.5  | 42.7 |
| $B_{\text{max},2}$\textsuperscript{b} | 29.7      | 10.3  | 11.0 |
| $n_1$\textsuperscript{c} | 1.50      | 1.72  | 1.00 |
| $n_2$\textsuperscript{c} | 8.80      | 25.1  | 29.1 |

$^a$First ($K_{d1}$) and second ($K_{d2}$) dissociation constant.

$^b$Binding component maxima, in terms of fractional occupancy (mol Ca\textsuperscript{2+} / mol hCasq1)

$^c$First ($n_1$) and second ($n_2$) Hill coefficient
Table 2. X-Ray Data Collection and Refinement Statistics

|                         | Wild-type hCASQ1 | D244G (low Ca$^{2+}$) | D244G (high Ca$^{2+}$) | M87T hCASQ1 |
|-------------------------|------------------|------------------------|-------------------------|-------------|
| **Data collection**     |                  |                        |                         |             |
| Space group             | C222             | P21/2;2                 | P21                     | P21         |
| Cell dimensions         |                  |                        |                         |             |
| $a$, $b$, $c$ (Å)       | 59.170, 145.132  | 66.106, 82.815         | 91.179, 67.462          | 65.681, 68.553, 99.262 |
| $a$, $b$, $c$ (°)       | 90.0, 90.0, 90.0 | 90.0, 90.0, 90.0        | 90.0, 90.0, 90.0         | 90.0, 92.845, 90.0 |
| Resolution (Å)          | 43.89 - 2.08     | 44.72 - 3.315          | 45.30 - 1.97 (2.04)     | 49.57 - 2.03 |
| $R_{merge}$             | 7.4 (76.9)       | 6.7 (20.7)             | 8.6 (44.0)              | 5.1 (33.0)  |
| $<I>/σI$                | 17.3 (2.6)       | 31.4 (10.6)            | 10.0 (2.3)              | 19.0 (5.0)  |
| Completeness (%)        | 99.4 (98.8)      | 99.7 (97.4)            | 96.1 (90.5)             | 99.6 (96.2) |
| Redundancy              | 6.8              | 7.0                    | 3.4                     | 3.7         |
| **Refinement**          |                  |                        |                         |             |
| Resolution              | 43.89 - 2.08     | 45.274 - 3.32          | 45.311 - 1.97           | 49.57 - 2.03 |
| Unique reflections      | 28782 (2822)     | 7652 (723)             | 129973 (12177)          | 57095 (5636) |
| $R_{work}$ / $R_{free}$| 0.1893 / 0.2168  | 0.2488 / 0.2904        | 0.1754 / 0.1931         | 0.1757 / 0.2074 |
| Number of atoms         |                  |                        |                         |             |
| Macromolecules          | 2,825            | 2,682                  | 11,425                  | 5,693       |
| Ion                     | 4                | 2                      | 82                      | 27          |
| Ligand                  | 16               | 16                     | 0                       | 0           |
| Water molecules         | 233              | 0                      | 2554                    | 727         |
| $B$-factors             |                  |                        |                         |             |
| Protein                 | 40.28            | 95.29                  | 21.7                    | 33.18       |
| Ligand/ion              | 48.98            | 95.38                  | 33.17                   | 38.84       |
| Water                   | 44.66            | ---                    | 32.41                   | 37.49       |
| R.m.s deviations        |                  |                        |                         |             |
| Bond lengths (Å)        | 0.002            | 0.002                  | 0.015                   | 0.002       |
| Bond angles (°)         | 0.510            | 0.510                  | 0.764                   | 0.600       |
| Ramachandran            |                  |                        |                         |             |
| Favored                 | 99.14            | 97.40                  | 98.50                   | 98.30       |
| Outliers                | 0                | 0                      | 0                       | 0           |
| Clashscore              | 2.19             | 3.32                   | 2.93                    | 3.09        |
Figure 1

A

\begin{align*}
\text{A280} & \quad \text{monomer} \\
0.0 & \quad 0.5 \\
0.5 & \quad 1.0 \\
1.0 & \quad 1.5 \\
1.5 & \quad 2.0 \\
2.0 & \quad 2.5 \\
\end{align*}

monomer

---

D244G

---

M87T

---

B

\begin{align*}
\text{A280} & \quad \text{dimer} \\
0.0 & \quad 0.5 \\
0.5 & \quad 1.0 \\
1.0 & \quad 1.5 \\
1.5 & \quad 2.0 \\
2.0 & \quad 2.5 \\
\end{align*}

dimer

---

C

\begin{align*}
\text{A280} & \quad \text{tetramer} \\
0.0 & \quad 0.015 \\
0.01 & \quad 0.010 \\
0.005 & \quad 0.005 \\
0.000 & \quad 0.000 \\
\end{align*}

tetramer

---

Elution volume (mL)

6.0

6.4

6.8

---
Figure 2

The graph illustrates the absorbance at 350 nm as a function of calcium concentration (Ca^{2+} in mM) for different conditions:

- **Wild-type**
- **2 mg/mL M87T**
- **4 mg/mL M87T**
- **D244G**

The absorbance values are plotted against the calcium concentration, showing distinct curves for each condition.
Figure 3
Figure 5
Figure 6

A  
Asp244  Pro246  Gly244  Pro246  Glu251

B  
Met87  Thr87
Figure 7

Normalized chain mean ADPs

- Wild-type
- Low-Ca$^{2+}$ D244G
- High-Ca$^{2+}$ D244G

Domain 1

Pro160

Domain 2

Thr263

Domain 3

Residue

Normalized chain mean ADPs

0.0

1.0

1.5

2.0

2.5

3.0

3.5

4.0

4.5

5.0

30 60 90 120 150 180 210 240 270 300 330 360 390
Figure 8
Characterization of Two Human Skeletal Calsequestrin Mutants Implicated in Malignant Hyperthermia and Vacuolar Aggregates Myopathy
Kevin M. Lewis, Leslie Ann Ronish, Eduardo Ríos and ChulHee Kang

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