S100A1 Protein Does Not Compete with Calmodulin for Ryanodine Receptor Binding but Structurally Alters the Ryanodine Receptor-Calmodulin Complex*

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S100A1 has been suggested as a therapeutic agent to enhance myocyte Ca2+ cycling in heart failure, but its molecular mode of action is poorly understood. Using FRET, we tested the hypothesis that S100A1 directly competes with calmodulin (CaM) for binding to intact, functional ryanodine receptors type I (RyR1) and II (RyR2) from skeletal and cardiac muscle, respectively. Our FRET readout provides an index of acceptor-labeled CaM (FKBP (FK506-binding protein 12.6) fluorescence) binding near donor-labeled FKBP (FK506-binding protein 12.6) on the cytoplasmic domain of RyR in isolated sarcoplasmic reticulum vesicles. S100A1 (0.01–400 μM) partially inhibited FRET (i.e. CaM binding), with Ki > 10 μM, for both RyR1 and RyR2. The high [S100A1] required for partial effects on FRET indicates a lack of competition by S100A1 on CaM/RyR binding under normal physiological conditions. High-resolution analysis of time-resolved FRET detects two structural states of RyR-bound CaM, which respond to [Ca2+] and are isoform-specific. The distribution of these structural states was perturbed only by high micromolar [S100A1], which promoted a shift of bound CaM to a lower FRET orientation (without altering the amount of CaM bound to RyR). Thus, high micromolar S100A1 does alter the CaM/RyR interaction, without involving competition. Nevertheless, submicromolar S100A1 can alter RyR function, an effect that is influenced by both [Ca2+] and [CaM]. We conclude that CaM and S100A1 can concurrently bind to and functionally modulate RyR1 and RyR2, but this does not involve direct competition at the RyR CaM binding site.

In striated muscle, the ryanodine receptor (RyR) is the channel responsible for Ca2+ release from the major intracellular Ca2+ reservoir, the sarcoplasmic reticulum (SR). RyR activity is tightly regulated by endogenous small molecules and proteins that maintain healthy Ca2+ cycling pertaining to relaxation and contraction of cardiac cells and skeletal muscle fibers. Two important modulators of Ca2+ cycling are EF-hand Ca2+-binding proteins, calmodulin (CaM) and S100A1. CaM is well known to directly bind to the RyR isoforms in skeletal (RyR1) and cardiac (RyR2) muscle and modulate channel activity in vitro or in situ (1, 2). S100A1 has also been shown to interact with and functionally modulate RyR1 and RyR2 (3–5). As a modulator of RyR2 activity, CaM is widely considered to be an effective inhibitor of Ca2+ leak through RyR2 (6, 7), and S100A1 is thought to exert a similar effect. Moreover, S100A1-based gene therapy shows promising effectiveness to correct abnormal Ca2+ cycling in heart failure (8, 9). To optimally exploit the therapeutic potential of S100A1, it is important to understand the structural basis of its molecular mode of action on the two principal targets hypothesized to mediate its effect on intracellular Ca2+: the RyR2 channels and the sarco/endoplasmic reticulum Ca2+-ATPase-phospholamban complex (10) that is responsible for active SR Ca2+ uptake. The present study is primarily focused on the S100A1/RyR2 interaction.

Similar to CaM, the tertiary structure of S100A1 and its modulatory action toward RyRs are affected by Ca2+ binding. With [Ca2+]cyto <0.5 μM, cytoplasmic exposure to nanomolar S100A1 has been shown to increase RyR1 activity, whereas ≥7 μM Ca2+ abolishes this modulatory action (3, 11). The apo (Ca2+-free)-CaM similarly increases RyR1 activity at nanomolar Ca2+, but the Ca2+-bound CaM decreases activity at high micromolar Ca2+ (1, 2, 12).

The modulatory action of S100A1 on RyR2 appears different from RyR1. In single-channel electrophysiology studies, cytoplasmic exposure to micromolar S100A1 has been shown to decrease the RyR2 channel activity at submicromolar and high micromolar Ca2+ (13). This in vitro action is consistent with reported effects of S100A1 in cardiomyocytes, whereby the addition of 0.1 μM S100A1 to permeabilized myocytes decreases RyR leak in resting conditions, and pressure injection of 0.1 μM S100A1 into cardiomyocytes increases depolarization-induced Ca2+ transient amplitudes (5). Thus, S100A1 and CaM might appear to exert similar effects on the RyR2 channel activity because CaM is well characterized to decrease RyR2 activity at both submicromolar and high micromolar Ca2+ (14,
late RyR activity and Ca\(^{2+}\) cycling in myocytes (4).

Given the high sequence identity between the CaM binding domains of RyR1 and RyR2, the S100A1 binding sites have been inferred to also be conserved between RyR1 and RyR2 (13). Taken together, these results have led to the hypothesis that S100A1 competes with CaM for the same binding site in RyR1 and RyR2. However, most studies that suggest competition between S100A1 and CaM use micromolar S100A1 and CaM to demonstrate this behavior. Moreover, S100A1 has micromolar affinity for the RyR peptide (16), which is inconsistent with the nanomolar [S100A1] shown to maximally modulate RyR activity and Ca\(^{2+}\) cycling in myocytes (4).

Here, we used established functional and state-of-the-art structural analysis approaches (18–20) to directly test the hypothesis that S100A1 competes with CaM binding to RyR1 or RyR2 in SR vesicles isolated from porcine skeletal muscle or hearts, respectively (Fig. 1). Our FRET toolkit is uniquely capable of resolving RyR-specific readouts from SR samples, in which RyRs are known to only be a small fraction (~25%) of the S100A1 targets in resting cardiomyocytes (6). Results largely contradict the tested hypothesis and show that S100A1 and CaM can concurrently bind to and regulate RyR1 and RyR2.

Results

Effect of S100A1 and CaM on \(^{3}H\)Ryanodine Binding to Cardiac and Skeletal SR—Because we are testing the hypothesis of competition between S100A1 and CaM for the same RyR binding site, we carried out all studies using SR preparations treated to remove residual CaM (as described under “Experimental Procedures”), thus tightly controlling any potential interference of CaM with S100A1 binding. In agreement with previous reports (21, 22), purified WT-S100A1 consisted of a mixture of molecules that had either retained or lost the N-terminal methionine. In addition, the S100A1 molecules were largely in the reduced, monomeric form, as determined from using electrospray ionization mass spectrometry. Although residual CaM is highly variable, it has not been controlled in the few previous studies that have assessed the modulatory action of S100A1 on RyR1 (3) and RyR2 (4, 23) activity in isolated SR membranes, where it may have led to confounding results. To investigate whether CaM alters the functional S100A1/RyR interaction, we evaluated the effect of 0.01–100 \(\mu\)M S100A1 on \(^{3}H\)ryanodine binding to CaM-stripped skeletal or cardiac SR, in the presence and absence of saturating (800 nM) CaM. We carried out these studies at 30 nM and 30 \(\mu\)M free Ca\(^{2+}\), thus reflecting cytosolic conditions in relaxed and contracting muscle cells. Despite the capacity for Ca\(^{2+}\) binding, the addition of 100 \(\mu\)M S100A1 marginally altered nanomolar and micromolar free Ca\(^{2+}\) by <20%. In previous FRET studies, we have shown that 800 nM CaM saturates the CaM binding sites on RyR (18, 20). Therefore, by using saturating CaM in these \(^{3}H\)ryanodine binding assays, we gauged the functional effect of S100A1 on RyR that is devoid of or fully loaded with CaM. In accord with previous studies (1, 2, 12), CaM significantly increased \(^{3}H\)ryanodine binding to skeletal SR (RyR1) in nanomolar Ca\(^{2+}\) by 10 ± 1-fold but decreased it in micromolar Ca\(^{2+}\) by 72 ± 1% (Fig. 2A). Saturating CaM lowered \(^{3}H\)ryanodine binding to cardiac SR (RyR2) in both 30 nM and 30 \(\mu\)M Ca\(^{2+}\) by 15 ± 1 and 29 ± 1% (Fig. 2A), respectively, which is also consistent with previous studies (14, 15).

We tested the functional effect of S100A1 on RyR1 in skeletal SR vesicles. In 30 nM Ca\(^{2+}\) and in the absence of CaM, 0.1–10 \(\mu\)M S100A1 inhibited \(^{3}H\)ryanodine binding in a biphasic profile, with maximum inhibition at 0.1 \(\mu\)M S100A1. With saturating CaM, however, \(^{3}H\)ryanodine binding was further increased by S100A1, but this tendency became significant only at 100 \(\mu\)M S100A1 (Fig. 2A, left). Thus, in the absence of CaM, S100A1 has the opposite effect of CaM on RyR1, whereas S100A1 may synergize with saturating CaM to enhance RyR1 activation in nanomolar Ca\(^{2+}\).

In micromolar Ca\(^{2+}\) and in the absence of CaM, ≤1 \(\mu\)M S100A1 had statistically significant, but modest, inhibitory effects on RyR1 \(^{3}H\)ryanodine binding, whereas 10 and 100 \(\mu\)M S100A1 had robust inhibitory effects (Fig. 2A, right). With saturating CaM, however, S100A1 had no effect on \(^{3}H\)ryanodine binding (Fig. 2A, right).
CaM/RyR Binding Is Independent of S100A1

The functional effect of S100A1 on RyR2 differs from its effect on RyR1. In nanomolar Ca\(^{2+}\) and in the absence of CaM (Fig. 2B, left), up to 10 \(\mu M\) S100A1 had negligible effects on cardiac SR \([^{3}H]\)ryanodine binding, but a robust increase (59 \pm 5\%) was seen in 100 \(\mu M\) S100A1. In saturating CaM, however, a statistically significant enhancement of \([^{3}H]\)ryanodine binding was revealed for all tested S100A1 concentrations. In fact, the inhibitory effect of CaM on RyR2 was completely overcome at 1 \(\mu M\) S100A1 and became a net RyR2 activation at 100 \(\mu M\) S100A1 (Fig. 2B, left). This is consistent with previously reported \([^{3}H]\)ryanodine RyR2-binding results (4).

In micromolar Ca\(^{2+}\) and in the absence of CaM, we found inhibition of \([^{3}H]\)ryanodine binding to RyR2, which is statistically significant only at high [S100A1] (Fig. 2B, right; 10 and 100 \(\mu M\) S100A1). With saturating CaM, we found a biphasic effect, activation of \([^{3}H]\)ryanodine binding plateauing in 0.1–10 \(\mu M\) S100A1 (Fig. 2B, right). This, too, is consistent with previously reported \([^{3}H]\)ryanodine RyR2 binding results (4).

Overall, these results establish that the S100A1 used in this study modulates RyR in concentrations ranging from 0.01 to 100 \(\mu M\), which cover the range reported for striated muscle tissue (3, 24). Whether this modulation is via orthosteric competition with CaM is explored further in this report.

**S100A1 Competition of AF568-labeled CaM (A-CaM) Binding to SR**—Two studies previously reported that micromolar CaM competes with S100A1-linked Sepharose for binding to skeletal SR vesicles and *vice versa* (16, 17), and these findings have been the primary basis of the hypothesis that these two proteins compete for overlapping binding sites on RyR1 and RyR2. However, given that S100A1 has other targets found in native SR preparations, there is a strong need to determine whether competition is specifically for RyR1. Furthermore, it is important to determine whether this mechanism also applies to CaM and S100A1 binding to RyR2, particularly given the strong therapeutic potential of S100A1 for heart failure (10, 23, 25–27). There are few data indicative of S100A1 interacting at the CaM binding site on RyR2. It was indirectly indicated in a report (13) that an RyR2 mutation (L3591D) abolishes both CaM- and S100A1-mediated modulation of RyR2 in nanomolar Ca\(^{2+}\). However, this is not definitive evidence of a direct interaction. For comparison with previous binding studies (16, 17) and with experiments shown below that examine competition of S100A1 with RyR-CaM binding, we first quantitated the level of S100A1 competing with CaM binding to *all* of its SR-associated targets (as opposed to only RyR). We utilized 0.1 \(\mu M\) A-CaM, to quantify, via fluorescence intensity of AF568 probe, the level of CaM co-sedimented with skeletal or cardiac SR vesicles following incubation with 0–400 \(\mu M\) S100A1. As a control, we show that a 200-fold excess of unlabeled CaM largely competes with the A-CaM retained by SR vesicles from both skeletal and cardiac muscle (Fig. 3, triangles). This control indicates that A-CaM binds to CaM binding sites endogenous to SR membranes (i.e. to all CaM-binding proteins in SR). A-CaM bound to skeletal SR was also significantly reduced upon incubation with \(\geq 100 \mu M\) S100A1, but this effect was more pronounced at 30 \(\mu M\) Ca\(^{2+}\) than at 30 \(\mu M\) Ca\(^{2+}\) with 17 and 45\%, respectively, of A-CaM bound (Fig. 3A). Conversely, preincubation of S100A1 equally reduced A-CaM binding to cardiac SR at both nano- and micromolar Ca\(^{2+}\), by as high as 33 and 42\%, respectively (Fig. 3B).

Overall, results shown in Fig. 3 are consistent with a significant overlap between the S100A1 and CaM binding sites within SR samples, particularly in micromolar Ca\(^{2+}\) (for skeletal muscle), and therefore they are in agreement with results from the competition assays that used S100A1- or CaM-conjugated Sepharose to retain rabbit SR vesicles (16, 17). Next, we tested whether S100A1 and CaM bind at overlapping sites on RyR1 and RyR2.

**S100A1 Competition of A-CaM Binding to RyR**—To resolve CaM binding specifically to RyR (as opposed to all CaM binding sites within SR), we measured FRET from donor-labeled FKBP

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**FIGURE 2.** CaM alters the effect of S100A1 on \([^{3}H]\)ryanodine binding to RyR1 and RyR2. SR vesicles from skeletal or cardiac muscle were incubated with S100A1 (0–100 \(\mu M\)) and CaM (0 or 800 nm) at 30 nm or 30 \(\mu M\) Ca\(^{2+}\) in the presence of \([^{3}H]\)ryanodine. A, as a control, the functional effect of 800 nm CaM was tested on \([^{3}H]\)ryanodine binding to skeletal (left) and cardiac (right) SR vesicles in 30 nm (blue) or 30 \(\mu M\) red) Ca\(^{2+}\). Data are shown as individual data points (square) and mean \(\pm\) S.E. (open circle). The fraction of \([^{3}H]\)ryanodine binding is relative to no CaM control. B, effect of S100A1 on \([^{3}H]\)ryanodine binding to skeletal SR (top panels) and cardiac SR (bottom panels) at 30 nm Ca\(^{2+}\) (left) and 30 \(\mu M\) Ca\(^{2+}\) (right) and in the absence of CaM (hollow squares) and in the presence of 800 nm CaM (solid squares). Control levels (absence of S100A1) are indicated by dashed lines. The fraction of \([^{3}H]\)ryanodine bound to SR vesicles is relative to maximum binding capacity (\(B_{\text{max}}\)). Data are shown as mean \(\pm\) S.E. \((n = 6–8)\), *, significant differences relative to no-S100A1 controls; \(p < 0.05\), as determined by paired Student’s \(t\)-test.
(D-FKBP) to A-CaM, with the donor and acceptor labels attached at residue 49 and 34, respectively. FRET between D-FKBP and A-CaM was measured as a decrease in donor fluorescence intensity, as we have shown previously (18–20, 28). We have previously established that this D-FKBP variant remains >90% bound to RyR over the time course of the experiment (18), which makes it ideal for providing an accurate and precise FRET readout. Furthermore, we have shown that these labeled FKBP and CaM variants have similar functional effects as WT-FKBP12.6 and WT-CaM on [3H]ryanodine binding to skeletal SR vesicles (18, 19). In the representative fluorescence spectra (Fig. 4, A and B) of skeletal SR samples labeled with D-FKBP, the decrease in fluorescence intensity due to FRET in the presence of 0.1 μM A-CaM was prevented by incubation with 20 μM unlabeled WT-CaM. By comparison, a much larger S100A1 concentration (10 μM) was required to elicit a decrease in the apparent A-CaM binding to RyR1. It is important to note that A-CaM is competed by WT-CaM with a K_i (∼0.1 μM) that is equivalent to the [A-CaM] used in this assay (Fig. 4), indicating that A-CaM and WT-CaM bind to the RyR1 with similar affinities. As shown in Fig. 4A, FRET to A-CaM in skeletal SR membranes is slightly reduced by preincubation of >30 μM S100A1 in micromolar, but not in nanomolar, Ca^{2+}. A reduction in FRET could be attributed to reduced binding or to a structural change within the D-FKBP-A-CaM-RyR1 macromolecular complex. However, the reduction in FRET that we observed is marginal and inconsistent with inhibition of CaM binding via the direct competition for the CaM binding site hypothesized based on studies using heavy skeletal SR (16, 17). Because these previous reports used SR from rabbit muscle, as opposed to the porcine SR used here, we tested the hypothesis of a species-dependent S100A1/CaM/RyR1 interaction by undertaking similar FRET experiments using rabbit skeletal SR and found that the effect of S100A1 on FRET is conserved between rabbit and pig skeletal SR (i.e. there is no competition between S100A1 and CaM for RyR1 binding) (data not shown). Using cardiac SR, we observed that ≥30 μM S100A1 fractionally decreases FRET between D-FKBP and A-CaM, and this effect is larger at nanomolar Ca^{2+} relative to micromolar Ca^{2+} (Fig. 4D). Given that FRET is subject to both the level of A-CaM bound to the RyR and the distance between FRET probes, the
S100A1-mediated reduction in FRET may be attributed to a reduction in CaM association with RyR relative to FKBP and/or to a structural change affecting the donor-acceptor distance. Experiments discussed further under "TR-FRET Analysis of the S100A1 Effects on RyR-CaM Structure and Binding" were designed to discern between these two mechanisms.

**CaM Competition of A-S100A1 Binding to RyRs**—To quantify S100A1 binding specifically to RyR, we measured FRET from D-FKBP to A-S100A1. To prepare A-S100A1, we used the WT single Cys (Cys-85) and also engineered several single-Cys S100A1 mutants, with the labeling site at T6C, T42C, and at the N and C termini via Ala-Cys-Ala tags at either end of S100A1. In the single-Cys S100A1 mutants, we substituted the single native Cys of WT-S100A1 (Cys-85) with either Ala (C85A) or Ser (C85S), as indicated (Fig. 5). In testing A-S100A1 labeled at its C terminus (AF568-S100A1-C85S-ACA), we were pleased to observe substantial FRET from D-FKBP to saturating [A-S100A1] in both RyR1 and RyR2 samples at 30 μM Ca²⁺ (Fig. 5, A and B). In 30 nM Ca²⁺, however, we observed insignificant FRET in the RyR1 sample and modest FRET in the RyR2 sample (Fig. 5, A and B). Saturation curves indicate A-S100A1/RyR binding with micromolar affinity. To our surprise, however, FRET was not perturbed by the addition of a molar excess of WT-S100A1, which indicates that this A-S100A1 construct is unlikely to be representative of WT-S100A1 binding. Furthermore, despite the lack of competition with WT-S100A1, FRET was fully abolished by the addition of the molar excess of WT-CaM. Moreover, we observed this trend (of selective competition of A-S100A1 by WT-CaM but not by WT-S100A1) for all labeled single-Cys S100A1 constructs tested (six different A-S100A1 constructs, as indicated in Fig. 5C). Explicitly, FRET from D-FKBP to nearly saturating A-S100A1 (2 μM), on both RyR1 and RyR2, was abolished by 20 μM WT-CaM, partially reduced by 20 μM unlabeled corresponding single-Cys S100A1 construct, and unaffected by 20 μM WT-S100A1 (Fig. 5, C and D). Given that our FRET readout, in this case, reflects binding of S100A1 to RyR in the proximity of D-FKBP, the lack of FRET inhibition by the addition of WT-S100A1 indicates that our A-S100A1 constructs do not reflect WT-S100A1 binding to both RyR1 and RyR2. Consequently, we did not pursue further studies with A-S100A1 in this project.

**TR-FRET Analysis of the S100A1 Effects on RyR-CaM Structure and Binding**—The D-FKBP to A-CaM FRET changes due to the addition of Ca²⁺ and/or S100A1 (Fig. 4) could be due to changes in either CaM binding or structure of the RyR-CaM complex. To resolve this ambiguity, we used TR-FRET, which allows simultaneous detection of both RyR-CaM binding and structure. The acceptor was attached at either CaM residue 34 (AF568 attached via CaM N-lobe; AN-CaM) or 110 (AF647 attached via CaM C-lobe; AC-CaM) of the full-length CaM, and the concentration of these acceptor-labeled CaMs was adjusted to give maximal FRET (18, 20). We tested the effect of 200 μM S100A1 because this concentration significantly altered FRET between D-FKBP and A-CaM in the competition experiments above (Fig. 4).

TR-FRET data were first analyzed to determine the fraction of donors participating in FRET (i.e. the fraction of D-FKBPs that are near an occupied CaM binding site). In all TR-FRET measurements, the fraction of uncoupled donors (i.e. donors not participating in FRET) was 10 ± 1%, and the addition of 200 μM S100A1 did not perturb this value, thus confirming that

![FIGURE 5. Competition of A-S100A1 by unlabeled S100A1 and CaM for binding to RyR. RyRs in SR membranes isolated from skeletal or cardiac muscle were labeled with D-FKBP and incubated with A-S100A1. Binding specificity of A-S100A1 to RyR was investigated by the addition of a >10-fold molar excess of unlabeled S100A1 or CaM. Saturation binding of AF568-S100A1-AF647-ACA to RyR from skeletal (A) and cardiac (B) SR is indicated as Total (solid lines) at 30 μM Ca²⁺ (blue) and 30 μM Ca²⁺ (red). Total FRET is compared with the saturation binding curve in the presence of 40 μM WT-S100A1 (dotted lines) or 40 μM WT-CaM (dashed lines). Binding specificities of several A-S100A1 variants to RyR from skeletal SR (C) or cardiac SR (D) were tested by the addition of 20 μM CaM (blue circles), WT-S100A1 (pink circles), or unlabeled S100A1 (green circles). Data are expressed as means ± S.E. (n = 4). *p < 0.05, as determined by analysis of variance with Fisher's post hoc test.]
CaM/RyR Binding Is Independent of S100A1

S100A1 does not compete with D-FKBP12.6 binding to RyRs. The 10% fraction of uncoupled donor is in accord with the previously reported slow D-FKBP/RyR dissociation rate (18).

Another advantage of TR-FRET is its ability to directly resolve multiple distances (i.e., structural states) within a protein complex as well as the population of probes in each structural state (29, 30). With AN-CaM, the TR-FRET data corresponding to both RyR isoforms (Fig. 6, A and B) fit best to a two-state Gaussian distribution model, characterized by a two-distance (R₁ and R₂) Gaussian distribution model of the separation between D-FKBP and AN-CaM within RyR. E and F, molar fractions of the populations associated with each of the two structural states characterized by distances R₁ and R₂. The S100A1 effects are shown in hatched bars, * significantly different from the no-S100A1 control. †, significantly different from 30 nM Ca²⁺, p < 0.05, as determined by analysis of variance with Fisher’s post hoc test.

Discussion

Using FRET methods that specifically resolve CaM binding to RyRs, we have tested the hypothesis that S100A1 modulates RyR activity via an interaction with the CaM binding site. This hypothesis has emerged based on the apparent competition of micromolar CaM or S100A1 with retention of RyR1 (in isolated skeletal SR vesicles) on Sepharose derivatized with either S100A1 or CaM, respectively (16, 17). Further supporting the tested hypothesis, S100A1 was shown to bind to an isolated 12-mer peptide that corresponds to RyR1 residues 3616–3627, which is a subsection of the 3614–3643 CaM binding domain 2 (16, 17), and mutation of one residue in this site (L3625D in RyR2) can abolish binding of S100A1 to the RyR peptide (11) and RyR channel modulation by S100A1 (11, 13). However, none of these studies have demonstrated that S100A1 binds specifically at the CaM binding site of full-length RyR. Indeed, semiquantitative analysis in a recent report indicated that increased expression of S100A1 in rat cardiomyocytes does not perturb CaM/RyR association (6). Our D-FKBP exclusively bind to RyRs in skeletal and cardiac SR
vesicles and, via FRET, they enable quantitative detection of A-CaM binding (within ±100 Å of D-FKBP) to intact, functional RyR1 or RyR2 in native SR membranes (18–20). Using this FRET system, we determined the structural influence of S100A1 on the CaM interaction with RyR1 and RyR2. To observe the reverse relationship (effect of CaM on the S100A1 binding sites but overlap with the CaM readout. However, all A-S100A1 binding sites differ from the endogenous S100A1 binding sites but overlap with the CaM binding site (Fig. 5, C and D). In addition to FRET-based quantification of CaM/RyR binding, we also used [3H]ryanodine binding to native SR membranes to determine whether CaM modulates the functional effect of S100A1 on RyR function (i.e. whether CaM and S100A1 may coexist within the RyR complex).

**S100A1 Marginally Affects CaM Binding to RyR1 and RyR2—** We found that FRET from D-FKBP to subsaturating A-CaMs is modestly reduced by a large (300–4000-fold) excess of S100A1 (less disordered) in micromolar Ca2+ (using AC-CaM). R3 is similar in RyR1 and RyR2 and is significantly shortened by Ca2+, but S100A1 does not affect this state (Table 1). Taken together, these results indicate that the N-lobe of CaM bound to RyR probes (peaks between) two structural states while anchored by a C-lobe in a single state. Measurable disorder, as indicated by FWHM (Table 1), is detected in all three distance measurements. Between RyR1 and RyR2, these states show significant, albeit small, differences both in their structural response to Ca2+ and to S100A1. However, TR-FRET data analysis detected no effect of S100A1 on the fraction of CaM associated with RyR1 or RyR2, as indicated by the unchanged fraction of uncoupled donor.

These results were surprising because it is thought that S100A1 modulates RyR function via the CaM binding site (16, 17). Several factors could explain this discrepancy between these and previous results. For the most part, high concentrations of Sepharose-linked S100A1 or CaM could have retained RyR1 indirectly due to its interaction with other SR membrane proteins. We show evidence of this in micromolar Ca2+ with >30 μM S100A1 reducing CaM co-sedimentation with skeletal and cardiac SR membrane proteins (Fig. 3). An additional or alternative explanation, perhaps even more likely, for the competition observed by Wright et al. (17) is indicated by the behavior of A-S100A1 illustrated in Fig. 5, where several different A-S100A1 constructs are competed from RyRs by WT-CaM not by WT-S100A1. This indicates that covalently modifying S100A1 (e.g. by attaching it to Sepharose, as done in Ref. 17) may change its binding specificity (relative to WT, unmodified S100A1), enabling the S100A1-Sepharose to attach to the CaM binding site of RyR1 or a site that is greatly impacted by CaM binding to RyR, rather than the true S100A1 binding site. This surprising behavior was replicated with several labeling sites, including Cys-85, Thr-6, Thr-42, and each terminus, thus suggesting that the specificity of S100A1 for a particular binding site on RyR is prone to change in response to covalent modification.

![Image](https://via.placeholder.com/150)

**CaM/RyR Binding Is Independent of S100A1**

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</tr>
<tr>
<td></td>
<td></td>
<td>200</td>
<td>48 ± 0.2</td>
<td>17 ± 2.7&lt;sup&gt;a,b&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup> Significant effect of Ca2+ (p < 0.05), as determined by analysis of variance with Fisher’s post hoc test.

<sup>b</sup> Significant effect of S100A1 (p < 0.05), as determined by analysis of variance with Fisher’s post hoc test.
12-mer peptide (included in the main CaM binding site) that is hypothesized to represent their binding site on RyR (16). For this application, the difference in FRET efficiency with A-CaM bound to RyR1 ($E \sim 0.3$) versus RyR2 ($E \sim 0.16$) is intriguing and is reminiscent of the isoform differences suggested via cryo-EM analysis in an elegant previous report (33).

In light of the marginal influence of S100A1 on CaM binding to RyR, it is surprising that a single-residue mutation within the CaM binding domain of RyR1 (L3625D) and RyR2 (L3591D) abolished their S100A1-mediated functional modulation (11, 13). Our results suggest that residues within the RyR CaM binding domain affect the S100A1/RyR interaction through an allosteric mechanism rather than through orthosteric, direct physical contact with S100A1.

**CaM Governs the Modulatory Effect of S100A1 on RyR Function**—We show that nanomolar S100A1 modestly but significantly modulates RyR1 and RyR2 function (Fig. 2B), despite our FRET results demonstrating that nanomolar S100A1 does not significantly alter the structural basis of CaM/RyR binding (Fig. 4). This indicates that nanomolar S100A1 does not functionally modulate RyR activity via the CaM binding site. Moreover, CaM does affect the S100A1-mediated functional regulation of RyR1 and RyR2, albeit in slightly different ways.

Our functional results (Fig. 2) suggest that there are at least two S100A1 binding sites on RyR1 at low Ca$^{2+}$: a nanomolar affinity inhibitory binding site that might be blocked by CaM/RyR binding and a micromolar affinity activatory binding site that is apparent when the inhibitory site is blocked by the presence of CaM. It also appears that S100A1 only mimics the action of CaM at micromolar Ca$^{2+}$, not nanomolar Ca$^{2+}$, indicating that S100A1 may bind to RyR1 at a site that overlaps with the CaM binding site in micromolar Ca$^{2+}$, but not in nanomolar Ca$^{2+}$. Given that S100A1 altered the conformation of the N-lobe, but not the C-lobe, of CaM binding to RyR (Table 1), it is plausible that micromolar S100A1 interferes with the CaM N-lobe interaction with RyR1 without significantly affecting the overall CaM/RyR1 apparent affinity, as previously speculated (34).

A report using saponin-skinned murine skeletal muscle fibers reported a dose-dependent increase in Ca$^{2+}$-induced SR Ca$^{2+}$ release when human recombinant S100A1 protein was added in the range from 0.01–1 μM, whereas 10 μM S100A1 tended to decrease isometric contractile force again (35). The inhibitory action of S100A1 is concurrent with our own observations of S100A1 reducing $[^3H]$ryanodine binding to RyR1, albeit with submicromolar S100A1 (Fig. 2B). This is noticeable in both nanomolar and micromolar Ca$^{2+}$ with CaM absent, which suggests that S100A1 may compensate for an absence of CaM in a skeletal muscle fiber, although the integration of signals from various accessory proteins probably contributes to the increased Ca$^{2+}$-induced SR Ca$^{2+}$ release observed by Most et al. (35).

Before this study, there was a general appreciation for the modulatory effect of S100A1 on RyR, although there were notable discrepancies in the literature, which probably reflect the complex and multifaceted interplay of RyR regulators. In 10 nM Ca$^{2+}$, the addition of submicromolar S100A1 has been shown to increase $[^3H]$ryanodine binding to skeletal SR membranes (RyR1), although this effect was abolished at 100 μM Ca$^{2+}$ (3). As mentioned, we demonstrate that submicromolar S100A1 decreases $[^3H]$ryanodine binding to RyR1, but this is restricted to the absence of CaM. Our data indicate (Fig. 2B) that this apparent discrepancy could be partially accounted for by residual endogenous CaM. Additionally, unaccounted impact of other accessory proteins as well as variation in the post-translationally modified state of RyR1 may also be responsible for the apparent discrepancies between studies.

At nanomolar Ca$^{2+}$, $[^3H]$ryanodine binding to RyR2 has been previously shown to be increased by 0.1 μM S100A1 but decreased by $>1$ μM S100A1 (4), whereas Yamaguchi et al. (13) have shown an inhibitory effect of submicromolar S100A1 on recombinant RyR2 in lipid bilayers. A possible cause for the discrepancy between reports and with our own data presented in this study is, again, the variable amount of residual endogenous CaM in SR membrane preparations, which we show greatly impacts the functional effect of S100A1 (Fig. 2).

Our results suggest that CaM binding to RyR2 enables an S100A1-mediated activation of RyR2 at both nanomolar and micromolar Ca$^{2+}$, which can override the inhibitory effect of CaM. This activation could be attributable to S100A1 competing with the inhibitory effect of CaM or to an allosteric interaction of CaM on S100A1-mediated modulation of RyR2. Given that micromolar S100A1 only modestly alters the CaM binding conformation, it appears more likely that the latter is occurring. Either way, these results require both S100A1 and CaM to concurrently bind to RyR2. Numerous studies in electrically stimulated and quiescent cardiomyocytes have indicated that S100A1 indeed has a dual effect on the activity of the macro-molecular RyR2 complex by improving both systolic opening and diastolic closure of the channel when intracellular S100A1 protein levels are elevated above normal (4, 6, 25). Increasing the amount of S100A1 bound to the RyR2 significantly reduced the calcium spark rate in normal cardiomyocytes and rescued the arrhythmogenic diastolic RyR2 dysfunction in failing cells. As such, these studies in intact cardiomyocytes partly resemble our experimental findings for $[^3H]$ryanodine binding gleaned from cardiac SR vesicle preparations when applied [S100A1] exceeded [CaM]. Overall, our results support the notion of S100A1 action being integrated in vivo within a complex context of modulatory signals from accessory molecules.

**Functional and Structural Effect of S100A1 under Diastolic and Systolic Ca$^{2+}$**—Our results indicate that the RyR1-S100A1 functional and structural interactions are more Ca$^{2+}$-sensitive than RyR2-S100A1 (Fig. 6 and Table 1). Based on the reported low Ca$^{2+}$ affinity of S100A1 ($K_a$ approximately low millimolar) (36), S100A1 is likely to remain in the apo-state in the concentrations tested. Thus, it is likely that the Ca$^{2+}$ dependence of the S100A1 structural effect on CaM-RyR1 is due to changes within RyR rather than S100A1. However, it is also possible that upon binding to RyR, the Ca$^{2+}$ affinity of S100A1 is drastically enhanced, as shown for CaM when it binds to RyR1 (37).

Post-translational modification of the only Cys of S100A1 (residue 85) has been shown to enhance Ca$^{2+}$ affinity to micromolar, which would facilitate the transition from apo- to Ca-S100A1 states. However, post-translational modifications are unlikely to be involved in our results, given that our mass...
spectrometry analyses indicated that S100A1 was unmodified before experiments, and the experimental conditions included the reducing agents GSH and DTT.

Possible S100A1 Binding Site(s) on RyR—Our results indicate a new model of S100A1/RyR interaction that involves S100A1 binding to more than one site on RyR and affecting the CaM-RyR structure but not the fractional occupancy of CaM binding sites on RyR. This is congruent with a report by Treves et al. (3) that proposed three S100A1 binding sites on RyR, including residues in RyR regions 1861–2155, 3774–3874, and 4425–4621. This encourages reassessment of these regions as candidates of S100A1 binding sites on RyR.

Despite some indications of competition between CaM and S100A1 binding to SR membranes and RyR, it should be understood that this is only observed when [S100A1] is in large (~1000-fold) molar excess. This is not entirely physiologically relevant when one considers a study indicating that the protein level of S100A1 is 9-fold greater than CaM in human hearts (38), with total [CaM] around 6 μM in the myocyte (39).

It may be interpreted from the lack of competition with CaM binding that S100A1 does not bind to RyR in our SR preparations. Despite not demonstrating a clear direct physical interaction between WT-S100A1 and RyR in our FRET-based assays, we clearly demonstrate that [3H]ryanodine binding is influenced by nanomolar [CaM], which thus indicates that S100A1 can functionally modulate RyR (Fig. 2A). In addition, further evidence of a physical interaction between RyR2 and S100A1 has been recently provided via proximity ligation assays and immunoprecipitation (6).

Conclusion—Although the hypothesis that CaM and S100A1 compete for the same RyR binding site is consistent with previous biochemical studies, our structural analysis of the RyR-CaM complex using TR-FRET reveals that S100A1 does not directly interfere with CaM binding to RyRs (Fig. 4) in the concentration range where CaM-dependent functional effects of S100A1 are observed (Fig. 2). Instead, S100A1 perturbs the structure of the RyR-CaM complex (Fig. 6). Thus, the structural mechanism for the interplay between CaM and S100A1 in regulation of RyR is more complex than previously thought.

Experimental Procedures

Isolation of SR Vesicles—SR membrane vesicles were isolated from porcine longissimus dorsi muscle and porcine cardiac ventricle by differential centrifugation of homogenized tissue, in accordance with Ref. 14. Heavy SR, rich in RyR1, was isolated by fractionation of skeletal crude SR vesicles using a discontinuous sucrose gradient (14). All vesicles were flash-frozen and stored at −80 °C. Immediately before the fluorescence binding studies described below, the SR vesicles were stripped of residual endogenous CaM by incubation with myosin light chain kinase-derived CaM binding peptide, followed by sedimentation, in accordance with Ref. 12.

Expression, Purification, and Fluorescence Labeling of FKBP and CaM—Single-cysteine mutants of FKBP12.6 (C22A/R49C/C76I, termed FKBP) and CaM (T34C or T110C) were separately expressed in *Escherichia coli* BL21(DE3)pLysS (Agilent Technologies), purified, and labeled with fluorescent probes Alexa Fluor 488 C5 maleimide (AF488), Alexa Fluor 568 C5 maleimide (AF568), or Alexa Fluor 647 C2 maleimide (AF647) (Life Technologies, Inc.), as described previously (19, 37). All fluorescently labeled proteins were diallyzed against a solution containing 30 mM NaCl and 20 mM MOPS (pH 7) and stored at −80 °C. Stoichiometric labeling of fluorescent CaM and FKBP to ≥95% was determined from the absorbance of bound dye relative to protein concentration determined via a bicinchoninic acid assay (Thermo Fisher Scientific) and SDS-PAGE densitometry. Essentially complete labeling was confirmed by electrospray ionization mass spectrometry.

Expression, Purification, and Fluorescence Labeling of S100A1—Single-cysteine variants (wild type and mutants: T6C/C85S, T6C/C85A, T42C/C85S, ACA-S100A1-C85S, and S100A1-C85S-ACA) of human S100A1 were expressed in *E. coli* BL21(DE3) and purified using an adapted protocol (40). A pRSETb T7 expression vector plasmid inserted with human S100A1 cDNA was transformed into *E. coli* BL21(DE3) pLysS, and the cells were grown at 37 °C until optical density at 600 nm reached ~1. At that point, protein expression was induced by the addition of 0.4 mM isopropyl-β-D-thiogalactoside and a further 3–4 h of incubation. The cells were spun, resuspended in a homogenization solution (2 mM EDTA, 1 mM EGTA, 1 mM TCEP, 1 mM PMFS, 1 μg/ml aprotinin/leupeptin, 1 mM benzamidine, 0.1% Tween 20, 31 μg/ml DNase I, 5 mM MgCl₂, and 50 mM Tris–HCl, pH 7.5), and briefly sonicated on ice with a Branson Ultrasonics Sonifier 250 (Danbury, CT). The homogenate was spun at 10,000 × g for 30 min. The supernatant was incubated for 20 min at 4 °C with 50% saturated ammonium sulfate. Following centrifugation, the ammonium sulfate saturation was increased to 80%, and the supernatant was incubated for 20 min at 4 °C and then centrifuged. S100A1 in the supernatant was precipitated by decreasing the pH to 4.1 and then pelleted by centrifugation. The pellet was resuspended in a solution containing 300 mM NaCl and 100 mM MOPS (pH 7) and dialyzed against 30 mM NaCl and 10 mM MOPS (pH 7). The S100A1 was further purified by loading onto a column containing phenyl-Sepharose CL-4B (GE Healthcare Bio-Sciences AB, Uppsala, Sweden) with 10 mM CaCl₂ and eluting using elution solution (500 mM NaCl, 5 mM EGTA, and 50 mM Tris pH 7.5). S100A1 was dialyzed against a solution containing 30 mM NaCl and 20 mM MOPS (pH 7.5) and concentrated using an Amicon stirred cell with a 3-kDa ultrafiltration disc (EMD Millipore, Billerica, MA). Protein purity and concentration were determined from SDS-PAGE and a bicinchoninic acid assay (Thermo Fisher Scientific), respectively. S100A1 was identified by electrospray ionization mass spectrometry using a QSTAR XL (AB Sciex, Framingham MA). In agreement with previous reports (21, 22), the WT-S100A1 sample contained a mixture of molecules that had either retained or lost the N-terminal methionine. In addition, the S100A1 molecules were largely in the reduced, monomeric form. A small fraction of protein contained the initiator N-formylmethionine, which has been observed previously (22).

Single-Cys variants of S100A1 were labeled with AF568 and purified using the same protocol as described previously for labeling of CaM (22). All fluorescently labeled S100A1 variants were dialyzed against a solution containing 30 mM
NaCl and 20 mM MOPS (pH 7) and stored at \(-80^\circ\)C. Stoichiometric labeling of fluorescent S100A1 to \(\geq95\%\) was determined from the absorbance of bound dye relative to protein concentration determined via a bicinchoninic acid assay (Thermo Fisher Scientific) and SDS-PAGE densitometry. Essentially complete labeling was confirmed by electrospray ionization mass spectrometry.

\[^{3}\text{H}]\text{Ryanodine Binding to SR Vesicles—}\] Skeletal and cardiac SR membranes (1 and 3 mg/ml, respectively) were preincubated with the indicated range of [S100A1], with and without 800 nM CaM, for 30 min at 4 \(^\circ\)C in a solution containing 150 mM KCl, 5 mM GSH, 1 \(\mu\)g/ml aprotinin/leupeptin, 1 mM DTT, 1 mM EGTA, 65 \(\mu\)M CaCl_2 (30 nM free Ca, as determined by MaxChelator), or 1.02 \(\mu\)M CaCl_2 (30 \(\mu\)M free calcium, as determined by MaxChelator), 0.1 mg/ml of BSA, and 20 mM K-PIPES (pH 7.0). Media containing S100A1 (100 \(\mu\)M) had a slightly shifted (<20%) free Ca\(^{2+}\), relative to the no-S100A1 controls, as determined using the Fura-2 calcium-sensitive dye or a calcium-sensitive electrode (Thermo Scientific, Beverly, MA) in solutions calculated for both nanomolar and micromolar free Ca\(^{2+}\), respectively. The assay solution at 30 \(\mathrm{nm}\) Ca\(^{2+}\) additionally contained 5 mM Na_2ATP, and 5 mM caffeine was also added to cardiac SR assays. Binding of \[^{3}\text{H}]\text{ryanodine (10 or 15 nM)}\) was determined following a 3-h incubation at 37 \(^\circ\)C and filtration through grade GF/B Glass Microfiber filters (Brandel Inc., Gaithersburg, MD) using a Brandel Harvester. In 4 ml of Ecolite scintillation mixture (MP Biomedicals, Solon, OH), the \(^{3}\text{H}\) retained on the filter was measured using a Beckman LS6000 scintillation counter.

Fluorescently Labeled CaM Binding to SR Vesicles—Skeletal SR (0.5 mg/ml) or cardiac SR (1 mg/ml) membranes were preincubated with the indicated [WT-CaM] or [S100A1] for 120 min, at 22 \(^\circ\)C in binding media (150 mM KCl, 5 mM GSH, 1 \(\mu\)g/ml aprotinin/leupeptin, 1 mM EGTA, 0.1 mg/ml BSA, and 20 mM K-PIPES, pH 7.0) supplemented with 65 \(\mu\)M CaCl_2 to give 30 \(\mathrm{nm}\) free Ca or with 1.338 mM CaCl_2 to give 300 \(\mu\)M free Ca\(^{2+}\) (as determined by MaxChelator). The binding of A-CaM to the treated SR membranes was measured following a 30-min incubation at 22 \(^\circ\)C with 0.2 \(\mu\)M A-CaM, centrifugation at 100,000 \(\times\) g for 25 min, and resuspension in a solution containing 5% SDS, 50 mM NaCl, 1 mM EGTA, and 20 mM Na-PIPES (pH 7.0). Bound A-CaM was determined from the fluorescence intensity at 600 nm (560-nm excitation, using a 570-nm emission long pass filter) (18).

FRET Measurements—Skeletal or cardiac SR (0.5 mg/ml) membranes were preincubated with 60 \(\mathrm{nm}\) AF488- or AF568-labeled FKBP (donor, D-FKBP), for 90 min, at 37 \(^\circ\)C, in a solution containing 150 mM KCl, 5 mM GSH, 0.1 mg/ml BSA, 1 \(\mu\)g/ml aprotinin/leupeptin, 1 mM DTT, and 20 mM PIPES (pH 7.0). To remove unbound D-FKBP, the SR membranes were spun at 110,000 \(\times\) g for 20 min and then resuspended to 3 mg/ml. These samples were then incubated with the indicated range of [WT-CaM] or [WT-S100A1] for 120 min at 22 \(^\circ\)C in binding media containing 30 \(\mathrm{nm}\) free Ca\(^{2+}\) or 1.02 \(\mu\)M CaCl_2 to give 30 \(\mu\)M free Ca\(^{2+}\) (calculated by MaxChelator). The binding of AF568- or AF647-labeled CaM (acceptor, A-CaM) or AF568-labeled S100A1 (A-S100A1) to RyR in the proximity of D-FKBP was measured following a 30-min incubation at 22 \(^\circ\)C with 0.1 \(\mu\)M A-CaM or 2 \(\mu\)M A-S100A1, unless otherwise indicated. Steady-state fluorescence spectra were acquired from samples in a 384-well, optical bottom, black wall plate using a Gemini EM microplate fluorometer (with a 490-nm excitation wavelength and a 495-nm long pass filter in the emission channel). FRET efficiency was measured as the fractional decrease of donor fluorescence intensity due to the presence of acceptor fluorophore, according to Ref. 19.

For TR-FRET measurements, we used time-correlated single-photon counting (41) or direct waveform recording, as described previously (29, 30). Global multiexponential analysis of the TR-FRET data was used to test a series of structural models, as described extensively elsewhere (30, 41). This mode of analysis resolves both binding and structural information from TR-FRET data. SR membranes labeled with D-FKBP were incubated with 200 \(\mu\)M S100A1 for 2 h at 22 \(^\circ\)C before a 30-min incubation at 22 \(^\circ\)C with 800 \(\mu\)M A-CaM.

Statistics—Sample means are from four or more independent experiments, and numbers of observations (n) are given in the figure legends. Each experiment was carried out using at least two independent SR preparations, isolated from different animals. Average data are provided as mean \(\pm\) S.E. Statistical significance was evaluated by use of either paired or unpaired Student’s t test or analysis of variance with Fisher’s post hoc test, as appropriate.

Author Contributions—R. T. R., D. D. T., D. M. B., and R. L. C. designed the study and wrote the paper. F. R. N. purified fluorescent proteins and undertook steady-state FRET experiments with cardiac SR membranes. D. R. and P. M. provided purified recombinant S100A1 protein and DNA, assisted with critical evaluation of the results, and edited the manuscript. R. T. R. purified unlabeled protein; labeled S100A1; and performed time-resolved and steady-state FRET studies, sedimentation assays, and \[^{3}\text{H}]\text{ryanodine binding assays using skeletal and cardiac SR membranes. All authors critically evaluated and approved the final version of the manuscript.}

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