Accessibility of Targeted DHPR Sites to Streptavidin and Functional Effects of Binding on EC Coupling

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In skeletal muscle, the dihydropyridine receptor (DHPR) in the plasma membrane (PM) serves as a Ca\(^{2+}\) channel and as the voltage sensor for excitation–contraction (EC coupling), triggering Ca\(^{2+}\) release via the type 1 ryanodine receptor (RyR1) in the sarcoplasmic reticulum (SR) membrane. In addition to being functionally linked, these two proteins are also structurally linked to one another, but the identity of these links remains unknown. As an approach to address this issue, we have expressed DHPR α\(_{1\alpha}\) or β\(_{1\alpha}\) subunits, with a biotin acceptor domain fused to targeted sites, in myotubes null for the corresponding, endogenous DHPR subunit. After saponin permeabilization, the ∼60-kD streptavidin molecule had access to the β\(_{1\alpha}\) N and C termini and to the α\(_{1\alpha}\) N terminus and proximal II–III loop (residues 671–686). Steptavidin also had access to these sites after injection into living myotubes. However, sites of the α\(_{1\alpha}\) C terminus were either inaccessible or conditionally accessible in saponin-permeabilized myotubes, suggesting that these C-terminal regions may exist in conformations that are occluded by other proteins in PM/SR junction (e.g., RyR1). The binding of injected streptavidin to the β\(_{1\alpha}\) N or C terminus, or to the α\(_{1\alpha}\) N terminus, had no effect on electrically evoked contractions. By contrast, binding of streptavidin to the proximal α\(_{1\alpha}\) II–III loop abolished such contractions, without affecting agonist-induced Ca\(^{2+}\) release via RyR1. Moreover, the block of EC coupling did not appear to result from global distortion of the DHPR and supports the hypothesis that conformational changes of the α\(_{1\alpha}\) II–III loop are necessary for EC coupling in skeletal muscle.

INTRODUCTION

In skeletal muscle, bidirectional signaling occurs between the dihydropyridine receptor (DHPR; a voltage-gated calcium channel composed of a pore-forming α\(_{1\alpha}\) subunit and auxiliary subunits α\(_{2}\), β\(_{1\alpha}\), and γ), and the ryanodine receptor (RyR1; a calcium release channel). Depolarization of the plasma membrane, where DHPRs are located, causes transmission of an orthograde signal from the DHPRs to the RyRs in the SR, resulting in calcium release (Rios and Brum, 1987; Tanabe et al., 1987; Garcia et al., 1994; Dirksen and Beam, 1999). This orthograde, excitation–contraction (EC) coupling signal is independent of the entrance of extracellular calcium. In addition, retrograde signaling exists whereby the association with RyR1 increases the magnitude of the voltage-gated Ca\(^{2+}\) current carried per DHPR (Nakai et al., 1996).

In addition to functional coupling of the DHPR and RyR1, structural coupling of these two proteins has been suggested from freeze-fracture studies of the plasma membrane, which reveal that DHPRs occur in “tetrads,” groups of four intramembranous particles that are arranged in ordered arrays (Franzini-Armstrong and Kish, 1995; Beam and Franzini-Armstrong, 1997; Protasi, 2002). The individual DHPRs within a tetrad are located in exact correspondence to the four subunits of RyR1 (Franzini-Armstrong and Kish, 1995; Block et al., 1988). The arrangement of DHPRs into tetrads is dependent on the presence of RyR1 (Protasi et al., 1998, 2000), implying that the DHPR and RyR1 are linked.

To understand skeletal-type EC coupling it is essential (a) to identify the regions of the DHPR that link it, directly or indirectly, to RyR1, and (b) to establish which of these regions may undergo voltage-driven conformational changes that are necessary for propagating the EC coupling signal. Toward this end, one approach has been to study cDNAs expressed in myotubes. Such approaches have shown that the α\(_{1\alpha}\) II–III loop critical domain (Nakai et al., 1998; Kugler et al., 2004) and the β\(_{1\alpha}\) C terminus are important for skeletal-type EC coupling (Beurg et al., 1999; Ahern et al., 2001; Sheridan et al., 2003), and that the α\(_{1\alpha}\) C terminus is important for targeting DHPRs to plasma membrane/SR junctions (Flucher et al., 2000b; Proenza et al., 2000). Additionally, in vitro biochemical studies have revealed that RyR1 has the ability to bind fragments of the α\(_{1\alpha}\) II–III loop and C terminus (Proenza et al., 2002), as well as the β\(_{1\alpha}\) C terminus (Cheng et al., 2005). However, considerable uncertainty remains as to whether these binding interactions also occur in vivo. Moreover, none of the studies to date have been able to distinguish whether the identified regions are involved in

Abbreviations used in this paper: BAD, biotin acceptor domain; 4-CMC, 4-chloro-m-cresol; DHPR, 1,4 dihydropyridine receptor; EC, excitation–contraction; FRET, fluorescence resonance energy transfer; RyR, ryanodine receptor; SA, streptavidin.
static interactions or undergo dynamic rearrangements during EC coupling.

As a new approach, we have begun to use cDNA constructs encoding a consensus sequence for metabolic biotinylation fused to sites of the DHPR. Previously, we employed this approach to demonstrate that after fixation and Triton permeabilization of myotubes expressing these constructs, the ∼60-kD molecule streptavidin has access to many sites of DHPRs that are inserted into fully assembled, plasma membrane/SR junctions, but that α₁S C-terminal regions may be occluded by RyR1 (Lorenzon et al. 2004).

The goal of the present work was to extend these studies to nonfixed myotubes, both to determine the pattern of streptavidin accessibility under conditions closer to those in vivo and to determine whether the binding of streptavidin interferes with the function of the DHPR as calcium channel and voltage sensor for EC coupling. We have found that in nonfixed, saponin-permeabilized cells, streptavidin has access to all those DHPR sites previously found to be accessible in fixed cells. However, α₁S C-terminal sites display a pattern of increased accessibility in nonfixed cells, suggesting that the C terminus exists in at least two conformations, one of which is closely apposed to RyR1. When injected into living myotubes, streptavidin was able to bind to the N and C termini of β₁s and to the N terminus of α₁S, but this binding did not affect electrically evoked contractions. By contrast, electrically evoked contractions were abolished when injected streptavidin bound to the proximal portion of the α₁S II–III loop, consistent with a model in which depolarization causes a rearrangement of the II–III loop such that the critical domain activates RyR1. These data represent the most direct evidence to date that a conformational change of the α₁S II–III loop is important for EC coupling.

MATERIALS AND METHODS

cDNA Constructs
The experiments described here made use of cDNA constructs (Fig. 1) that contained sequence for a biotin acceptor domain (BAD) of either 70 or 97 amino acids. The BAD sequence was designated YFP-BAD (previously described (Tanabe et al., 1988) in a single nucleus with a cDNA construct (5–100 ng/l). After injection, the cells were washed into a medium in which the biotin present in the 2% horse serum was supplemented with an additional 1 μM biotin.

Expression of cDNA
Primary cultures of myotubes isolated from newborn dysgenic (αᵢₛ-null) or β₁-null mice were prepared as described previously (Beam and Franzini-Armstrong, 1997). Myoblasts were plated on ECL-coated (Upstate Biotechnology) glass-bottom (MatTek), or primaria plastic (BD Biosciences) 35-mm culture dishes. Myotubes were grown for 6–7 d in a humidified 37°C incubator with 5% CO₂. Approximately 1 wk after plating, myotubes were microinjected (Tanabe et al., 1988) in a single nucleus with a cDNA construct (5–100 ng/μl). After injection, the cells were changed into a medium in which the biotin present in the 2% horse serum was supplemented with an additional 1 μM biotin.

NeutrAvidin Staining and Imaging
2 d after injection, myotubes were washed with an “internal” solution containing (in mM) 140 Cs-aspartate, 10 Cs₂-EGTA, 5 MgCl₂, and 10 HEPES, pH to 7.4 with CsOH. The myotubes were permeabilized with saponin (12 μg/ml in internal solution) for 30 s, and then exposed for 30 min in the dark to a 1:2,000 dilution (in internal solution) of NeutrAvidin-tetramethylrhodamine (Molecular Probes), hereafter referred to as ”streptavidin.” The cells were washed twice with internal solution, fixed with 4% paraformaldehyde in PBS for 20 min, and then imaged with an Axiovert/LSM 510 META laser-scanning confocal microscope (Carl Zeiss Microimaging, Inc.). Excitation/emission parameters for each fluorophore were as follows: YFP, 488 nm excitation, 488/543 nm dual dichroic, 505–530 nm band-pass emission filter; rhodamine, 543 nm excitation, 488/543 nm dual dichroic, 560 nm long-pass emission filter. Cells were viewed with 40× (1.3 NA) or 63× (1.4 NA) oil immersion objectives.

Evoked Contractions and Streptavidin Injection
Myotubes were bathed in a rodent Ringers solution containing (in mM) 146 NaCl, 5 KCl, 1 MgCl₂, 10 HEPES, 11 glucose (pH to 7.4 with NaOH) and tested for electrically evoked contractions (extracellular stimulation 100 V, 10–20 ms). Cells that produced robust contractions were then bathed in Ringers solution containing N-benzyl-β-toluenesulphonamide (BTS; 10 μM) for 5–10 min to block contractions during subsequent microinjection with streptavidin (1 mg/ml). The cells were rinsed twice with culture medium and returned to the tissue culture incubator for at least 2 h. The identified myotubes were then retested for evoked contractions under the same conditions as before streptavidin injection. Records of evoked contractions were obtained by measuring the movement of an identifiable portion of a myotube across the visual field from confocal images acquired at a rate of 11 Hz.

Calcium Transients and Streptavidin Injection
To analyze intracellular Ca²⁺ transients in response to membrane depolarization or to direct ryanodine receptor activation, intact myotubes were loaded with the fluorescent calcium indicator dye Fluo-3 AM. Before imaging, myotubes were washed free of the culture medium with rodent Ringer and exposed to the dye (5 μM in rodent Ringer) for 1 h at 37°C. Calcium transients were recorded in response to membrane depolarization by 80 mM KCl or RyR1 activation by 0.5 mM 4-chloro-m-cresol (4-Cmc), which were applied for 5 s via an extracellular pipette. Calcium transients were measured as a change in fluorescence site at the distal α₁₅ C terminus (residue 1860) and introduce a new KpnI site after residue 1667 (forward primer GCCAATGCA-CAAGGTACCTATGGCAACGAGCAGCATAG; reverse primer ATCCGGGCGGGCGGTAGCGTGTGAGCAGGTCCT). The plasmids were cut with KpnI, and then the appropriate plasmid fragments were ligated to produce YFP-α₁₅(1667-BAD)1800. Restriction digests and sequencing were used to verify the construct.

Streptavidin Binding to Biotinylated DHPR Sites

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intensity ($\Delta F/F$) using confocal imaging (excitation/emission parameters for fluo3-AM dye: 488 nm excitation, 505 nm LP emission filter).

Whole-Cell Recording Methods
Calcium currents were recorded with the whole-cell variant of the patch clamp technique. Pipettes were fabricated from borosilicate glass and had resistances of 1.5–2.4 MΩ when filled with internal solution (composition given above). For the measurement of membrane currents, the external solution contained (in mM): 145 tetraethylammonium-Cl, 10 CaCl$_2$, 10 HEPES, and 0.003 tetrodotoxin, pH 7.4 with tetraethylammonium-OH. Test currents were corrected for linear components of leak and capacitative currents by digital scaling and subtraction of averaged control currents elicited by 20-mV hyperpolarizing steps from the holding potential (~80 mV). Cell capacitance was determined by integration of these same control currents and used to normalize test currents (pA/pF). To inactivate T-type currents and isolate L-type currents, a 1-s prepulse to $-30$ mV followed by a 100-ms repolarization to $-50$ mV was applied before the test pulse.

RESULTS

Fig. 1 illustrates the constructs used, all of which contained BAD at one site and fluorescent protein at a second site as an independent reporter of DHPR localization. Except for YFP-$\alpha_{1S}(1667$-BAD)$1860$, these constructs were examined in a previous study (Lorenzon et al., 2004) and found able to restore electrically evoked contractions. We also found this to be the case for YFP-$\alpha_{1S}(1667$-BAD)$1860$ expressed in $\alpha_{1S}$-null myotubes (12/13 cells). Thus, the incorporation of a BAD did not appear to interfere with the targeting or essential function of the $\alpha_{1S}$ or $\beta_{1a}$ fusion proteins.

Streptavidin Accessibility of Targeted DHPR Sites

Our previous work (Lorenzon et al. 2004) showed that in fixed and Triton-permeabilized myotubes, streptavidin had access to all of the sites illustrated in Fig. 1, except for BAD fused to the end of either the proximal (YFP-$\alpha_{1S}$(1667-BAD)-$\beta_{1a}$) or distal (YFP-$\alpha_{1S}$(1860-BAD)) C terminus. Both of these sites were occluded in the presence of RyR1. An important goal of the present studies was to assay streptavidin accessibility under conditions similar to those in vivo. As one part of this effort,

Figure 1. Schematic illustration of the BAD-DHPR fusion constructs. Constructs are designated by the sites of attachment of BAD (red circle) and YFP (yellow oval). In the $\alpha_{1S}$ II–III loop constructs, $\alpha_{1S}$ residues 672–685 were replaced by the BAD. For the $\alpha_{1S}$ C terminus, BAD was attached after residue 1860, or attached after residue 1667, or placed after 1667 followed by $\alpha_{1S}$ 1668–1860 (YFP-$\alpha_{1S}$(1667-BAD)$1860$).

Figure 2. In nonfixed myotubes, the $\alpha_{1S}$ N terminus, the $\alpha_{1S}$ proximal II–III loop, and the N and C terminus of $\beta_{1a}$ are accessible to streptavidin (A–D, respectively). Left panels illustrate the localization of the YFP-DHPR fusion proteins (indicated in green), center panels illustrate the localization of the bound streptavidin (red), and right panels show the overlay. Bars, 5 μm.
nonfixed cells expressing the DHPR-BAD fusion proteins were saponin permeabilized, and then exposed to streptavidin. Saponin-permeabilized myotubes retain substantial function as shown by both spontaneous Ca\(^{2+}\) release and caffeine-induced global Ca\(^{2+}\) release (Ward et al., 2000, 2001).

As in fixed myotubes, BAD fused to the α\(_{1S}\) N terminus was accessible in nonfixed myotubes to the ~60-kD streptavidin molecule (Fig. 2 A). Specifically, red fluorescent foci (center panel), representing bound streptavidin, were colocalized with the foci of fluorescent protein (indicated in green, left), which report the distribution of the α\(_{1S}\) subunit. BAD inserted into the proximal region of the α\(_{1S}\) II–III loop (replacing α\(_{1S}\) residues 671–686) was also accessible to streptavidin binding (Fig. 2 B) in nonfixed myotubes. A similar pattern of colocalizing red and green foci was present for the N and C termini of the β\(_{1S}\) subunit (Fig. 2, C and D). Thus, the accessibility of all these sites to streptavidin was similar in fixed and nonfixed myotubes.

The only DHPR site that showed a difference in streptavidin accessibility in nonfixed versus fixed myotubes was the proximal portion of the α\(_{1S}\) C terminus. In fixed dysgenic myotubes (Lorenzon et al., 2004), both the distal α\(_{1S}\) C terminus (YFP-α\(_{1S}\)1860-BAD) and proximal, truncated α\(_{1S}\) C terminus (YFP-α\(_{1S}\)1667-BAD) were inaccessible to streptavidin. In nonfixed cells, the distal α\(_{1S}\) C terminus (YFP-α\(_{1S}\)1860-BAD) was also inaccessible. The lack of accessibility is illustrated in Fig. 3 A, where YFP-α\(_{1S}\)1860-BAD and streptavidin puncta did not colocalize (especially apparent in the overlay image, right). However, in distinction to fixed myotubes, the proximal, truncated α\(_{1S}\) C terminus (YFP-α\(_{1S}\)1667-BAD) was at least partially accessible in nonfixed myotubes (see both the presence and absence of streptavidin puncta colocalized with the YFP puncta in Fig. 3 B). To investigate this partial accessibility further, we examined streptavidin binding to BAD inserted at the identical, proximal position, but followed by the additional, C-terminal sequence, YFP-α\(_{1S}\)(1667-BAD)1860. The presence of this additional C-terminal sequence caused BAD at residue 1667 to be accessible to streptavidin in nonfixed myotubes (Fig. 3 C).

Table I summarizes the extent of colocalization between streptavidin binding and clusters of DHPRs. The first data column presents the judgment of the authors, based on identified images. The majority of these same images were also randomly assembled and presented unidentified to four individuals who assigned each image a numerical value of 0 (<5% colocalization), 1 (5–95% colocalization), or 2 (>95% colocalization). The mean values for these scores are presented in Table I, data column 2, and agree well with the authors’ assessment of identified images.

**Table 1**

| DHPR/BAD construct | # cells exhibiting colocalization | Colocalization score |
|--------------------|----------------------------------|----------------------|
| BAD-β\(_{1S}\)-YFP  | 12 (12)                          | 1.9 (11)             |
| YFP-β\(_{1S}\)-BAD  | 16 (16)                          | 1.8 (11)             |
| BAD-α\(_{1S}\)-YFP  | 15 (13)                          | 1.8 (7)              |
| α\(_{1S}\)(671-BAD-686)-YFP | 12 (12)                          | 1.7 (10)             |
| YFP-α\(_{1S}\)1860-BAD | 1 (1)                            | 0.1 (12)             |
| YFP-α\(_{1S}\)1667-BAD | 30 (59)                          | 0.8 (17)             |
| YFP-α\(_{1S}\)(1667-BAD)1860 | 21 (21)                          | 1.9 (12)             |

The total number of cells examined for each construct noted in parentheses. The YFP-α\(_{1S}\)1860-BAD and YFP-α\(_{1S}\)1667-BAD cells exhibited partial colocalization (considerably fewer dots were colocalized compared to the other constructs).

Figure 3. Differential accessibility of α\(_{1S}\) C-terminal sites to streptavidin in nonfixed myotubes. (A) Myotubes expressing YFP-α\(_{1S}\)1860-BAD did not display binding of streptavidin that colocalized with loci of fluorescent DHPRs. (B) Myotubes expressing YFP-α\(_{1S}\)1667-BAD displayed incomplete colocalization between loci of DHPRs and streptavidin binding. The arrow indicates a cluster of DHPRs with substantial binding of streptavidin, and the arrowhead a cluster of DHPRs showing very little binding of streptavidin. (C) Nearly complete colocalization was observed for YFP-α\(_{1S}\)(1667-BAD)1860. Bars, 5 μm.
it was necessary first to release unbound streptavidin from the cell, which was accomplished by saponin permeabilization. Fig. 4 A2 shows for a normal myotube (which lacks DHPR-BAD fusion protein) that virtually no unbound streptavidin remains following this permeabilization. Fig. 4 (B and C) illustrates myotubes expressing either BAD-α1S-YFP or YFP-β1a-BAD, which were injected with streptavidin and 1 h later subjected to saponin permeabilization. The colocalization of red and green foci indicates that streptavidin binding had occurred during the previous 1-h incubation. As a final control experiment, normal myotubes were tested for EC coupling 2–5 h after injection of streptavidin. Extracellular stimulation of such myotubes elicited brief, strong contractions (29/29 cells; unpublished data).

To test the effects of streptavidin binding on EC coupling, individual dysgenic or β1-null myotubes expressing DHPR/BAD fusion constructs were initially tested for electrically evoked contractions before streptavidin injection (Fig. 5, top). At least 2 h after injection, the identified cells were retested for evoked contractions (Fig. 5, bottom). Evoked contractions were not affected by streptavidin binding to the N or C terminus of β1a (10/11 and 18/20 cells contracted, respectively) or to the α1S N terminus (10/11 cells). However, binding of streptavidin to the proximal portion of the α1S II–III loop abolished evoked contractions (Fig. 5, weak contraction in 1/12 cells; no contractions in the remainder). Thus, the II–III loop was the only DHPR site at which the additional mass of streptavidin interfered with EC coupling.

Electrically evoked contractions require that a brief depolarization produce substantial Ca2+ release. Thus, a partial reduction of Ca2+ release might account for the loss of electrically evoked contractions after streptavidin binding to the proximal II–III loop. To test this, we examined Ca2+ release in response to a more prolonged depolarization produced by application of 80 mM KCl. Fig. 6 (top) shows that Ca2+ released by a 5-s depolarization was reduced to a nearly undetectable level by streptavidin binding to the proximal II–III loop. This nearly complete suppression of EC coupling was not a consequence of depletion of SR Ca2+ stores or a direct effect of streptavidin on RyR1. Specifically, Fig. 6 (bottom) shows that Ca2+ release in response to the RyR agonist, 4-CmC, was not obviously different in streptavidin-injected, α1S(671-BAD-686)-YFP–expressing myotubes.

In principle, the binding of streptavidin to the α1S II–III loop could have prevented a specific conformational change necessary for EC coupling without affecting other functions of the DHPR. Alternatively, this binding could have induced a widespread distortion of α1S that abolished function of the DHPR more globally. Fig. 7 illustrates calcium currents in myotubes expressing α1S(671-BAD-686)-YFP that were injected

Figure 4. Diffusion and binding of streptavidin injected into living myotubes. (A1) 1 h after streptavidin (SA) injection of a normal myotube, diffuse red fluorescence could be observed throughout the cell. (A2) In the absence of DHPR-BAD fusion proteins, injected streptavidin is rapidly released after saponin permeabilization. 1 h after SA injection of a normal myotube, saponin was applied and an image obtained ~15 min later with laser intensity, detector/amplifier gains, and offset set to be the same as for the images in B. (B) A 1-h period was sufficient to allow binding of injected streptavidin to BAD-α1S-YFP expressed in a dysgenic myotube. Nonbound streptavidin was released by saponin permeabilization. Bars, 5 μm. (C) Colocalization of red and yellow fluorescent puncta from injected streptavidin-rhodamine and YFP-β1a-BAD in a β1-null myotube. Note that BAD-β1a-YFP streptavidin binding has not yet been confirmed by confocal imaging.

Functional Impact of Streptavidin Binding
The ability of streptavidin to bind at DHPR sites in nonfixed myotubes raises the question of whether the addition of this large mass interferes with the function of the DHPR as the voltage sensor for EC coupling. To address this question, saponin permeabilization was not an appropriate method of applying streptavidin, which was introduced instead by intracellular microinjection. For this approach, it was important to determine (a) whether injected streptavidin could diffuse and bind to DHPR sites within intact myotubes, and (b) whether injected streptavidin interferes with the function of normal myotubes. Fig. 4 A1 demonstrates that injected streptavidin (1 mg/ml) uniformly filled a large myotube within 1 h after injection. To determine if this period of time was also sufficient for binding,
with streptavidin. Streptavidin had no obvious effect on either the voltage dependence or amplitude of \( \text{Ca}^{2+} \) currents in these cells. The average peak amplitude of currents in \( \alpha_{1S}(671\text{-BAD}-686)\)-YFP–expressing cells was similar whether or not streptavidin was injected (Fig. 7 B) and comparable to that of dysgenic myotubes expressing \( \alpha_{1S}-YFP \) (Bannister and Beam, 2005). Thus, the ability of \( \alpha_{1S}(671\text{-BAD}-686)\)-YFP to function in EC coupling was eliminated by streptavidin binding, whereas its function as a calcium channel and its retrograde regulation from RyR1 were not affected. Although not systematically examined, we also found no obvious effect of injected streptavidin on \( \text{Ca}^{2+} \) currents in myotubes expressing \( \beta_{1a-BAD}, \beta_{1a-YFP} \), or \( \alpha_{1S}-YFP \) (unpublished data).

**DISCUSSION**

In this paper, we have examined the consequences of applying streptavidin to myotubes expressing DHPRs with targeted sites of biotinylation. Except for the \( \alpha_{1S} \) C terminus, all other DHPR sites examined, including the \( \alpha_{1S} \) N terminus and proximal II–III loop and the \( \beta_{1a} \) N and C termini, were accessible to streptavidin binding and showed equivalent accessibility in nonfixed myotubes as found previously (Lorenzon et al., 2004) for fixed cells. However, the accessibility of the \( \alpha_{1S} \) C terminus differed between fixed and nonfixed cells and depended on the location of the BAD, the length of the C terminus, and the presence or absence of RyR1. In addition to examining accessibility, we also characterized the functional consequences of streptavidin binding at sites of the \( \alpha_{1S} \) and \( \beta_{1a} \) subunits. Excitation–contraction coupling was abolished by streptavidin binding to the proximal portion of the \( \alpha_{1S} \) II–III loop, but was unaffected by binding at the \( \alpha_{1S} \) N terminus or \( \beta_{1a} \) N or C terminus.

**Streptavidin Accessibility**

Except for the \( \alpha_{1S} \) C terminus, all other sites of \( \alpha_{1S} \) and \( \beta_{1a} \) showed equivalent accessibility in fixed and nonfixed cells. The ability of streptavidin to bind to the C terminus appears to depend on the location of the BAD within the C terminus, the length of the C terminus,

**Figure 5.** Excitation-contraction coupling is abolished by streptavidin binding to the proximal portion of the \( \alpha_{1S} \) II-III loop, but is unaffected when binding occurs at the \( \alpha_{1S} \) N terminus or \( \beta_{1a} \) N or C terminus. Electrically evoked contractions are shown for intact \( \beta_{1a} \)-null myotubes expressing \( \text{YFP-}\beta_{1a-BAD} \) or \( \text{BAD-}\beta_{1a-YFP} \), or dysgenic myotubes expressing \( \text{BAD-}\alpha_{1S-YFP} \) or \( \alpha_{1S}(671\text{-BAD-686})\)-YFP. Prior to streptavidin injection, myotubes responded to extracellular stimulation with robust contractions (top). 2–4 h after streptavidin injection, the same individual myotubes were tested for their responses to identical stimulation (bottom). Vertical scale, arbitrary units. For \( \text{BAD-}\beta_{1a-YFP} \)-expressing cells, streptavidin binding following injection has not yet been confirmed by confocal imaging.

**Figure 6.** Effect of streptavidin binding to the proximal \( \alpha_{1S} \) II–III loop on depolarization- and 4-CmC–induced \( \text{Ca}^{2+} \) transients. Representative responses are shown for normal myotubes (A) and SA-injected \( \alpha_{1S}(671\text{-BAD-686})\)-YFP–expressing myotubes (B) to a 5-s application of either 80 mM KCl (top) or 0.5 mM 4-CmC (bottom). \( \Delta F/F \) scale bar = 0.4 or 0.6 for the KCl responses of the normal and SA-injected myotubes, respectively, and 1.0 or 0.6 for the 4-CmC responses of the normal and SA-injected myotubes, respectively. (C) Summary of peak \( \Delta F/F \) amplitudes for KCl (top) and 4-CmC (bottom) responses of either normal myotubes (Norm) or SA-injected \( \alpha_{1S}(671\text{-BAD-686})\)-YFP–expressing myotubes (SA-inj).
and the presence of RyR1. The distal C terminus (residue 1860) is not accessible in either fixed or nonfixed cells, suggesting that this region may be buried. However, the accessibility of the proximal C terminus (residue 1667) is more complex. The BAD is partially accessible when attached at the end of a truncated (after residue 1667) C terminus in nonfixed myotubes (Fig. 3 B), but inaccessible in fixed myotubes (Lorenzon et al., 2004). Our data do not allow us to differentiate between the possibilities that the proximal α1S C terminus is partially accessible in all junctions or ranges from accessible to inaccessible, perhaps as a consequence of developmental maturity of the junctions. In either case, it is necessary to suppose that fixation reduces the partial accessibility of the proximal α1S C terminus to below the level of detection.

The partial accessibility of the proximal α1S C terminus in nonfixed myotubes could also reflect the existence of two, functionally distinct conformations, one of which is occluded by RyR1 and one which is not. If the fractional occupancy of the nonoccluded state was only a few percent, nearly all sites would be occluded after fixation. Furthermore, the binding of streptavidin in nonfixed cells would gradually trap the proximal C terminus in the nonoccluded conformation, resulting in partial colocalization of streptavidin with junctional DHPRs. However, BAD attached at the identical, proximal site (1667) within a longer C terminus (BAD followed by α1S residues 1668–1860) was accessible to streptavidin in nonfixed myotubes (Fig. 3 C), suggesting that the distal C terminus may anchor the proximal C terminus in a nonoccluded conformation.

The pattern of streptavidin accessibility to DHPR sites is in good agreement with a previous study using the complementary approach of measuring the fluorescence resonance energy transfer (FRET) efficiency of a CFP-YFP tandem attached to these same sites of DHPRs within junctions that either did, or did not, contain RyR1 (Papadopoulos et al., 2004). In that FRET study, a difference in FRET was taken as an indication that the presence of RyR1 impacted the environment of the attachment site. In agreement with the present work, the
FRET efficiency was equivalent in the presence and absence of RyR1 when the reporter was attached to the α1S N terminus and proximal II–III loop, and to the β1a C terminus. The two methods were also in quite good agreement for the α1S C terminus truncated at 1667, where the presence of RyR1 resulted in incomplete accessibility to streptavidin and reduced CFP-YFP FRET. The β1a N terminus was the only site for which the two methods gave different results in that the presence of RyR1 did not prevent streptavidin accessibility but did alter CFP-YFP FRET. One could imagine that the β1a N terminus is close to RyR1 but that biotin projects from the attached BAD in an orientation that allows streptavidin accessibility.

The proximal C-terminal site examined in our studies (residue 1667) lies near the site at which naturally occurring, proteolytic cleavage has been reported to cause truncation of ~95% of α1S in adult skeletal muscle (De Jongh et al., 1991; Hulme et al., 2005). With respect to the distal site (residue 1860) examined in the present study, it is important to note that a much larger fraction (~30%) of α1S in cultured myotubes is full length (Shulman, 1996). Moreover, Catterall and colleagues (Hulme et al., 2005) propose that the A-kinase anchoring protein, AKAP15, anchors the cleaved portion of the C terminus to the proximal portion. In any case, there is good evidence that a large fraction of α1S in myotubes has the distal C terminus attached (Flucher et al., 2000a), although the nature of this attachment remains unclear.

A number of possible roles have been ascribed to the C-terminal tails of L-type Ca2+ channels. In particular, portions of the C terminus of α1C (Gao et al., 2000) and α1S (Proenza et al., 2000) may be important both for trafficking to the plasma membrane, and for targeting to plasma membrane junctions with the SR (Flucher et al., 2000b; Proenza et al., 2000). Moreover, biochemical studies have suggested that the C-terminal residues 1393–1527 of α1S bind calmodulin and can bind to RyR1 (Sencer et al., 2001). Additionally, both the α1S C terminus (residues 1431–1435) and RyR1 contain consensus binding sites for Homer (Xiao et al., 2000). Perhaps the direct or indirect linkage of α1S C-terminal residues to RyR1 has the consequence that the distal C terminus of α1S is inaccessible to streptavidin in RyR1-containing junctions, either because RyR1 directly occludes the distal C terminus or because the RyR1-induced arrangement of DHPRs into tetrad reds results in occlusion.

Functional Impact of Streptavidin Binding: Direct Evidence that EC Coupling Depends on a Conformational Change Involving the α1S II–III Loop

Previous work has implicated a variety of DHPR sites as important for EC coupling, including the II–III loop critical domain and the III–IV loop of α1S and the C terminus of the β1a. In particular, studies using expression in myotubes null for α1S or β1a have shown that mutations and/or deletions within the aforementioned regions profoundly affect EC coupling (Nakai et al., 1998; Beurg et al., 1999; Ahern et al., 2001; Sheridan et al., 2003; Kugler et al., 2004). A priori, these regions could be directly involved in protein–protein interactions linking DHPRs to RyRs, but they could equally well impact the ability of other regions of the DHPR to participate in such interactions. Additionally, current candidate regions of the DHPR could either be involved in static interactions that link DHPRs to RyRs, or actively undergo conformational changes that are essential for propagating the EC coupling signal. Unfortunately, the functional analyses to date do not provide a basis for distinguishing these possibilities.

Here we have found that excitation–contraction coupling is abolished by streptavidin binding to the proximal portion of the α1S II–III loop, but is unaffected when binding occurs at the α1S N terminus or β1a N or C terminus. Fig. 8 illustrates a model of EC coupling together with mechanisms by which streptavidin binding could interfere with this coupling. In particular, EC coupling is pictured (Fig. 8 A) as dependent upon a conformational change of the α1S II–III loop with the result that the critical domain (residues 720–765, pink) contacts RyR1 and activates it to release Ca2+ from the SR. Conceivably, the binding of streptavidin to the proximal II–III loop could distort the overall structure of the DHPR so as to entirely abolish the function of this protein (Fig. 8 B). This possibility would appear to be excluded by the result that calcium currents are unaffected by the binding of streptavidin to the II–III loop. Importantly, the amplitude of DHPR Ca2+ current depends not only on the structural integrity of the DHPR but also on a retrograde functional interaction between RyR1 and the DHPR (Nakai et al., 1996). This retrograde interaction is abolished by structural alterations of the critical domain of the α1S II–III loop (Grabner et al., 1999; Kugler et al., 2004). Thus, the fact that Ca2+ current amplitude was unaffected by streptavidin binding to the proximal II–III loop provides additional support that this binding does not significantly distort the spatial interrelationship between the DHPR and RyR1.

As an alternative to a global disruption of structure, bound streptavidin could specifically obstruct conformational changes of DHPR cytoplasmic domains required for EC coupling. As shown in Fig. 8 C, streptavidin bound to the proximal II–III loop interferes with the conformational change that causes the downstream, critical domain to contact RyR1. An alternative possibility is that the critical domain contacts RyR1 at rest and that bound streptavidin prevents a torsional conformational change that is necessary for activation of Ca2+ release. In any case, our results provide the most direct evidence to date that conformational changes of the α1S II–III loop are critical for EC coupling, but do not
preclude the possibility of additional, important conformational changes of other DHPR cytoplasmic domains.

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