$\alpha_2 \delta_1$ Dihydropyridine Receptor Subunit Is a Critical Element for Excitation-Coupled Calcium Entry but Not for Formation of Tetrads in Skeletal Myotubes

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ABSTRACT It has been shown that small interfering RNA (siRNA) partial knockdown of the $\alpha_2\delta_1$ dihydropyridine receptor subunits cause a significant increase in the rate of activation of the L-type Ca²⁺ current in myotubes but have little or no effect on skeletal excitation-contraction coupling. This study used permanent siRNA knockdown of $\alpha_2\delta_1$ to address two important unaddressed questions. First, does the $\alpha_2\delta_1$ subunit contribute to the size and/or spacing of tetradic particles? Second, is the $\alpha_2\delta_1$ subunit important for excitation-coupled calcium entry? We found that the size and spacing of tetradic particles is unaffected by siRNA knockdown of $\alpha_2\delta_1$, indicating that the visible particle represents the α_{1s} subunit. Strikingly, >97% knockdown of $\alpha_2\delta_1$ leads to a complete loss of excitation-coupled calcium entry during KCI depolarization and a more rapid decay of Ca²⁺ transients during bouts of repetitive electrical stimulation like those occurring during normal muscle activation in vivo. Thus, we conclude that the $\alpha_2\delta_1$ dihydropyridine receptor subunit is physiologically necessary for sustaining Ca²⁺ transients in response to prolonged depolarization or repeated trains of action potentials.

INTRODUCTION

The dihydropyridine receptor (DHPR) complex in the plasma membrane/transverse tubular system of skeletal muscle is composed of five subunits ($\alpha_1, \alpha_2, \beta, \delta$, and γ) and serves at least three distinct functions. One of these functions is as a high-voltage-activated, L-type Ca²⁺ channel. As for other voltage-activated Ca²⁺ channels, the structures underlying voltage sensing and ion permeation reside in the α_1 subunit (termed α_{1s} , CaV1.1 in skeletal muscle). In addition to this function as a calcium channel, the CaV1.1 serves to link depolarization of the plasma membrane to Ca²⁺ release from the type 1 ryanodine receptor (RyR1) in the sarcoplasmic reticulum (SR), which in turn causes contraction. This excitation-contraction (EC) coupling in skeletal muscle does not depend on the L-type Ca^{2+} current and is thought instead to reflect conformational coupling between the CaV1.1 and RyR1. Strong support for such conformational coupling is provided by the discovery of the retrograde interaction between the two molecules (1,2) and freeze fracture electron microscopy. The latter technique reveals that DHPR complexes are arranged in groups of four (termed "tetrads") such that each tetradic particle, resulting from the fracture of a

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single complex, is aligned with one of the four identical subunits of an underlying RyR1 (3).

A third function of the DHPR complex is in the recently elucidated process termed "excitation-coupled calcium entry" (ECCE). In ECCE, depolarization causes the entry of extracellular Ca²⁺ via a pathway with pharmacological similarities to the pathway involved in store-operated calcium entry (SOCE) (4,5). However, ECCE differs from SOCE in that it does not require the emptying of intracellular Ca²⁺ stores and it requires the presence of both the CaV1.1 and RyR1. Until our recent discovery (4,5), evidence for depolarization associated Ca²⁺ entry had only been indirectly demonstrated (6-8), but the mechanism for this entry remained elusive. In addition, the fact that muscle twitches (9) could be sustained in Ca²⁺-free external buffer focused attention away from seeking mechanisms for this entry, even though it had been known for a long time that prolonged contractions could not be sustained for long periods of time in the absence of external Ca^{2+} (10,11). Discrimination between depolarization induced SR Ca^{2+} release and sarcolemmal Ca²⁺ entry is technically difficult and therefore has limited investigations of ECCE in adult skeletal muscle fibers. However, recent results from studies using the Mn^{2+} quench technique on single dissociated flexor digitorum brevis fibers have recapitulated the existence of our ECCE results from myotube experiments; ECCE is seen in adult skeletal muscle fibers, and its rate is enhanced by ryanodine pretreatment (12).

An important tool for analyzing how the functions of the DHPR complex depend on its various subunits has been provided by animals bearing null mutations. Clearly important is

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CaV1.1, which consists of four repeat sequences, each having six transmembrane helices. EC coupling is absent in skeletal muscle of dysgenic mice, which are homozygous null for CaV1.1 (13–19), and there is an absence of large particles in the sarcolemma of these animals when subjected to freeze fracture studies (3). EC coupling is also absent in skeletal muscle of mice or zebrafish null for the cytoplasmic β_1 subunit (16–20). In both mice and zebrafish, the absence of the β_1 subunit results in an impaired trafficking of α_{1S} to the plasma membrane. Additionally, the work on zebrafish has shown that those α_{1S} subunits that do reach the plasma membrane lack a well-defined association with RyR1 in β_1 null fish (20). Genetic deletion of the γ subunit, which contains four transmembrane helices, does not markedly affect muscle function (13,21).

The α_2 and δ subunits contribute almost half of the total mass of the DHPR and arise by cleavage of a common precursor. The α_2 subunit is positioned on the extracellular side of the plasma membrane and is disulfide bonded to the δ subunit, which has a single transmembrane span (22). Mice null for the $\alpha_2 \delta$ isoform that is expressed in skeletal muscle $(\alpha_2 \delta_1)$ appear to have an embryonic or birth lethal phenotype (J. Offord, Pfizer, Groton, CT, personal communication, 2007). Thus small interfering RNA (siRNA) knockdown represents a useful alternative for examining the roles of $\alpha_2 \delta_1$ in skeletal muscle. Previously, it was shown that siRNA partial knockdown of $\alpha_2 \delta_1$ caused a significant increase in the rate of activation of the L-type Ca²⁺ current in myotubes but had little or no effect on skeletal EC coupling (23). The goal here was to use siRNA knockdown to address two important questions not examined in these previous studies. We chose the myotube system rather than adult fibers because only with myotubes is it possible to study a defined population of cells with the identical amount of gene silencing in each cell and because we previously demonstrated that the two systems recapitulate each other in areas we wished to study.

The first question posed was, is the $\alpha_2\delta_1$ DHPR subunit important for ECCE (4,5)? The second question was, does the $\alpha_2\delta_1$ subunit contribute to the size and/or spacing of tetradic particles? With respect to the latter question, we found that the size and spacing of tetradic particles is unaffected by siRNA knockdown of $\alpha_2\delta_1$, which may indicate that the $\alpha_2\delta_1$ protein does not affect the fracturing process and that the visible particle represents the α_{1s} subunit. Strikingly, in response to the first question we found that knockdown of $\alpha_2\delta_1$ leads to a nearly complete loss of physiologically stimulated ECCE, which results in a rapid decrement in the size of myoplasmic Ca²⁺ transients during bouts of repetitive stimulation like those occurring during normal muscle activation in vivo.

MATERIALS AND METHODS

Cell culture and siRNA transduction

Primary murine myoblasts were cultured on collagen-coated 10 cm plates (Costar, Cambridge, MA) for propagation and on Matrigel (BD Bioscience, San Jose, CA) coated plates (96-well plates (Opticlear Costar 3614) for imaging, and 10 cm dishes for membrane preparations) in Ham's F10 20% fetal bovine serum, 5 ng/ml basic fibroblast growth factor, 100 μ g/ml streptomycin sulfate, 100 units/ml penicillin-G in 5% CO₂. Myoblasts were transduced three times with one of two different pSiREN retrovirus constructs (Fig. 1 *A*) containing $\alpha_2\delta_1$ complementary sequences connected by a central hairpin loop SIR1: 5'... G TGT GAG CGG ATT GAC TCT ... 3', and SIR2: 5'... G GCA CGC TTT GTG GTG ACA ...3' (Fig. 1 *A*) and

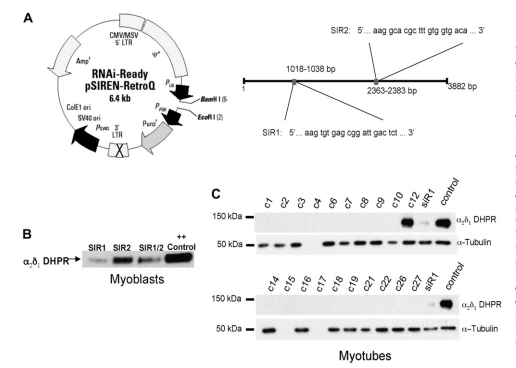


FIGURE 1 siRNA vector construction and effects of $\alpha_2 \delta_1$ siRNA knockdown on expression of $\alpha_2 \delta_1$ protein in primary myoblasts and myotubes. (A) Sequences of siR1 and siR2 and vector map of retroviral construct used to transduce primary muscle cells. (B) Western blot showing expression of $\alpha_2 \delta_1$ in pools of primary myoblasts after transduction three times with retrovirus expressing constructs for siR1 and siR2. (Note siR2 did not significantly lower $\alpha_2 \delta_1$ expression.) (C) Western blots showing expression of $\alpha_2 \delta_1$ in myotubes derived from 27 individual myoblast clones that had been transduced three times with retrovirus expressing siR1. α -tubulin (*lower band*) used as the loading control. All clones with the exception of C12 showed negligible expression compared to control. At the right extracts from siR1 pooled myoblasts and control myoblasts are shown as controls.

selected for puromycin resistance conferred by the vector. Resistant cells were then plated at clonal density and 20 individual clones screened for $\alpha_2\delta_1$ messenger RNA (mRNA) expression by TaqMan real time polymerase chain reaction (PCR) analysis to select for cells expressing <2% of control. A previously created TRPC3 siRNA knockdown cell line was used as a control for possible off-site nonspecific effects of siRNA (24). For Ca²⁺ imaging, electrophysiology, freeze fracture, or biochemistry transduced cells were allowed to reach 60%–70% confluence and were induced to differentiate into myotubes for 5 days by changing the growth medium to Dulbecco's modified Eagle's medium containing 2% heat-inactivated horse serum, 100 µg/ml streptomycin sulfate, and 100 µg/ml penicillin-G in 20% CO₂.

Membrane preparation and immunoblotting

Crude membrane preparations were made from myotubes after 5 days in differentiation media. Myotubes were harvested in harvest buffer (137 mM NaCl, 3 mM KCl, 8 mM Na₂HPO₄, 1.5 mM KH₂PO₄, and 0.6 mM EDTA, pH 7.2) from 10–15 10 cm plates and centrifuged for 10 min at $250 \times g$. The pellet was resuspended in buffer consisting of 250 mM sucrose, 10 mM HEPES, pH 7.4, supplemented with 1 mM EDTA, 10 µg/ml leupeptin, 0.7 μ g/ml pepstatin A, 5 μ g/ml aprotinin, and 0.1 mM phenylmethanesulfonyl fluoride and then homogenized using a Polytron cell disrupter (Brinkmann Instruments, Westbury, NY). The whole cell homogenates were centrifuged for 20 min at 1500 \times g max, and the supernatants were collected and recentrifuged for 60 min at 100,000 \times g max at 4°C. The membrane pellets were finally resuspended in 250 mM sucrose, 20 mM HEPES, pH 7.4, frozen in liquid N₂, and stored at -80° C. Western blot analysis was done on a heavy SR fraction from control and $\alpha_2 \delta_1$ DHPR knockdown myotubes using a method published previously (25). The membrane fractions were solubilized in sodium dodecyl sulfate and size fractionated by polyacrylamide gel electrophoresis. The polyacrylamide gels were then blotted onto polyvinylidene fluoride membranes and, after blocking, hybridized with the appropriate primary antibody. They were then probed with a horseradish peroxidase secondary antibody and level of expression determined with chemilumenescence. The samples were probed for RyR1, CaV1.1, $\alpha_2\delta_1$, skeletal calsequestrin (CSQ), FK506-binding protein 12 kDa (FKBP12), SR/ER Ca^{2+} -ATPase (SERCA1), and skeletal triadin (25). Anti- β -tubulin (26) was used as a loading control.

Quantitative PCR

TaqMan Real Time PCR analysis was performed on total RNA extracted from two 10 cm plates of differentiated myotubes using RNeasy Kit (Qiagen, Valencia, CA). $\alpha_2\delta_1$ mRNA levels relative to glyceraldehyde-3-phosphate dehydrogenase (GAPDH) mRNA levels were determined by quantitative reverse transcription-polymerase chain reaction using predeveloped commercial TaqMan Gene Expression Assays from Applied Biosystems (Assay ID: Mm00486607_m1; Foster City, CA) according to the manufacturer's instructions in an ABI PRISM 7700 Sequence Detection System for 50 cycles. $\alpha_2\delta_1$ /GAPDH mRNA ratios were obtained from the equation $2^{-\Delta CT}$, where ΔCT is the difference in threshold cycles between $\alpha_2\delta_1$ and GAPDH.

Immunohistochemistry analysis

Cells were fixed in cold methanol (27,28). After blocking, fixed cells were hybridized with an anti- $\alpha_2\delta_1$ primary mouse monoclonal antibody (20A, Hybridoma Studies Bank, University of Iowa) and then hybridized with a goat anti-mouse Cy3 labeled secondary antibody and observed with a 100× oil immersion lens on an Olympus IX70 microscope (Olympus, Tokyo, Japan).

Whole-cell measurements of Ca²⁺ currents

 Ca^{2+} currents were measured (29,30) using whole-cell patch-clamping with a Warner PC501 patch amplifier (Hamden, CT). Cells were patched with pipettes that were pulled from borosilicate glass capillaries to a resistance of 1.7–3.0 MΩ. Pipettes were filled with internal solution containing (in mM) 140 CsAsp, 5 MgCl₂, 10 Cs₂ EGTA, and 10 HEPES, with the pH adjusted to 7.4 with CsOH. After a seal was formed, the bath was perfused with an additional ~10 ml of the external solution containing (in mM) 145 tetra-ethylammonium (TEA), 10 CaCl₂, 0.003 tetrodotoxin, and 10 HEPES, pH 7.4, with CsOH. Control currents, elicited by 20–40 ms voltage steps from -80 mV to -110 mV, were used to determine linear capacitance (cells >800 pF were excluded from further analysis). The voltage clamp command sequence was to sequentially step from: a holding potential of -80 mV, to -20 mV (200 ms), to -50 mV (75 ms), to the test potential (20 ms), to -50 mV (125 ms), and back to -80 mV. Test potentials ranged in 10 mV increments from voltages from -20 to +60 mV/+80 mV. Test currents were corrected for linear components of leak and capacitive currents by digital scaling and subtraction of the average of 10 control currents, which were elicited with the pulse protocol just described, with a 200 ms test potential of -80 mV.

Peak current density-voltage relationships of individual cells were fitted with the equation

$$I = G_{\max} \times \frac{(V - V_{rev})}{1 + e^{-\frac{V - V_{1/2}}{k}}}$$

where *I* is the peak current density at test potential *V*, G_{max} the maximum Ca^{2+} conductance, V_{rev} the reversal potential, $V_{1/2}$ the potential for halfmaximal activation of conductance, and *k* the slope factor. Time constants of current activation were determined by fitting individual current records with a single ($\alpha_2 \delta_1$ knockdown) or double (normal) exponential function.

All data are presented as mean \pm SE.

Freeze fracture and measurements of intratetrad distance

The cells were washed twice in phosphate buffer saline at 37° C, fixed in 3.5% glutaraldehyde in 0.1 M sodium cacodylate buffer, pH 7.2, stored in fixative for 1–4 weeks, and briefly infiltrated in 30% glycerol. A small piece of the coverslip was mounted with the cells facing a droplet of 30% glycerol, 20% polyvinyl alcohol on a gold holder, and then frozen in liquid nitrogen-cooled propane. The coverslip was flipped off to produce a fracture that followed the cell surface originally facing it. The fractured surfaces were shadowed with platinum at 25° while rotating and then replicated with carbon in a Balzers freeze fracture machine (model BFA 400; Balzers Spa, Milan, Italy). Replicas were photographed in a Philips 410 Electron Microscope 410 (Philips Electron Optics, Mahwah, NJ). The specimen holder was positioned in the eccentric position to minimize variations in electron optical magnification. Parallel culture dishes were immunolabeled (see Immunohistochemistry analysis above).

Measurements of intratetrad distance

The distances between the pale center of each black ring of platinum (representing a DHPR complex) and the centers of its nearest neighbors in a tetrad were measured using the NIH Image software on micrographs taken at a magnification of 33,900 and scanned at a resolution of 1200 dpi. Measurements on $\alpha_2\delta_1$ knockdown cells were done on clones C12 and C14. To minimize variations due to distortion of the proteins during fracturing, measurements were taken only from portions of tetrads in which the angle subtending the lines connecting three adjacent particles was 90° or close to it. Incomplete tetrads with one particle missing as well as complete four-particle tetrads were used for the measurements. Blind data collection was insured by randomly mixing images from control and knockdown cells.

Myotube Ca²⁺ imaging

Stock concentrations of caffeine solutions were prepared in imaging buffer (125 mM NaCl, 5 mM KCl, 2 mM CaCl₂, 1.2 mM MgSO₄, 6 mM glucose, and 25 mM HEPES, pH 7.4). In KCl-containing solutions the concentration of NaCl was adjusted as necessary to maintain the total ionic strength (KCl + NaCl) equal to 130 mM.

Differentiated myotubes were loaded with 5 μ M Fluo-4 AM (Molecular Probes, Eugene, OR) at 37°C for 20 min in imaging buffer as described previously (31). The cells were then washed three times with imaging buffer and transferred to a Nikon TE2000 microscope (Tokyo, Japan). Fluo-4 was excited at 494 nm, and fluorescence emission was measured at 516 nm using a Nikon 40× FluorApo oil 1.3 numerical aperture (na) objective. Data were collected with an intensified 12-bit digital intensified charge-coupled device (ICCD; Stanford Photonics, Stanford, CA) from regions consisting of 10–20 individual cells and analyzed using QED software (QED, Pittsburgh, PA). Test agents were delivered using the Multivalve Perfusion System (Automate Scientific, Oakland, CA). A dose response curve for a single agent was performed by adding an ascending concentration of that agent in any given well to compare the response to any given agent of the cells expressing the $\alpha_2\delta_1$ siRNA to that of cells expressing wild-type (*wt*) $\alpha_2\delta_1$ protein.

To compare different experiments, individual response amplitudes were normalized to the maximum fluorescence obtained in the same cell depolarized with 60 mM KCl. To test for differences in Ca^{2+} stores, a group of both *wt* and $\alpha_2\delta_1$ knockdown cells were exposed to 200 nM thapsigargin (TG) in the presence of nominal Ca^{2+} -free media and the Ca^{2+} release response recorded. To test that a deficiency in Ca^{2+} entry during depolarization might be responsible for the observed responses of $\alpha_2\delta_1$ knockdown cells to depolarization, cells were pretreated with nominal Ca^{2+} -free buffer for 1 min before depolarization in the same nominal Ca^{2+} -free buffer with 60 mM KCl. Where indicated, nonlinear regression with sigmoidal dose-response analysis was performed using Prism version 4.0. (GraphPad Software, San Diego, CA). Data are presented as mean ± SE.

Additional experiments were performed to measure cell responses to electrical pulse trains. Myotubes were loaded with Fluo-4 as described above and fluorescence emission (516 nm) acquired with an IC-300 ICCD camera (Photon Technology International, Lawrenceville, NJ). Data were captured, digitized, and stored on computer using ImageMaster software (Photon Technology International). Bipolar electrical field stimuli were applied using two platinum electrodes fixed to opposite sides of the well and connected to an AMPI Master 8 stimulator set at 3 V, 25 ms pulse duration, over a range of frequencies (0.05–20 Hz). In these experiments, data were acquired at 200 ms intervals. In separate experiments, a fatigue stimulus protocol was applied. Electrical pulses (7 V, 1 ms duration) were applied at 100 Hz for 500 ms interspersed with 500 ms rest for a total duration of 5 min. The frequency-transient amplitude relationship was studied using 3 s pulse trains in a range of 1–120 Hz.

Mn²⁺ quench and electrical field stimulation

Mn²⁺ influx was measured as previously described (32,33). Briefly, differentiated primary *wt* or $\alpha_2\delta_1$ knockdown myotubes were loaded with fura-2, the cells were imaged using an ICCD camera with an Olympus 40× oil 1.2 na objective, and data were collected from 3–10 individual cells at five frames per second. The isosbestic wavelength for fura-2 was calibrated at the beginning of each experiment. A final concentration of 500 μ M MnCl₂ (in the presence of 1.2 mM external Mg²⁺) was used, and emission was monitored at 510 nm. Cells loaded with fura-2 (5 μ M) were stimulated with either local application of K⁺ or by electrical field stimulation as described above. To compare the effects of $\alpha_2\delta_1$ DHPR knockdown on both traditional store depletion induced entry (SOCE) and ECCE, myotubes were pretreated for 10 min with 200 nM TG followed by confirmation that the stores were empty with a challenge with 20 mM caffeine. Mn²⁺ quench experiments were only performed on cells that did not respond to the caffeine challenge (>99%).

RESULTS

Construction of siRNA vectors that efficiently knock down expression of the $\alpha_2 \delta_1$ DHPR subunit

We used the following general design guidelines to construct our two siRNA vectors: 1), find 21 nt sequences in the target mRNA that begins with an adenine dinucleotide, 2), select several target sequences with 30%–60% GC content and no stretches of >4 Ts or As at different positions along the length of the gene sequence, and 3), compare the potential target sites to the appropriate genome database to eliminate from consideration any target sequences with more than 16–17 contiguous basepairs of homology to other coding sequences.

Fig. 1 A shows a map of the location of the two selected siRNA sequences in the $\alpha_2 \delta_1$ complementary DNA (cDNA). Because myoblasts are difficult to transfect at high efficiency and because of the need to have multiple copies of the vector permanently integrated in the cell to reduce expression of the target mRNA to <2% of control, a pSiREN retrovirus was used to transduce the cells at moiety of infection = 5 on three separate occasions (34,35). Between transductions, cells were selected with puromycin to eliminate any cells that were not transduced with at least one copy of the vector from the cell population. Western blot analysis of pooled cells after the third round of infection with the siRNA retrovirus demonstrated that although both siR1 and 2 reduced the expression of $\alpha_2 \delta_1$ in myoblasts, siR1 was much more efficient at this task (Fig. 1 B). Cells from pool 1 were plated at clonal density and the clones expanded. As can be seen in Fig. 1 C, myotubes derived from most of the clones from this pool did not produce sufficient $\alpha_2 \delta_1$ protein to be detected by Western blot analysis.

In Fig. 2 A, TaqMan analysis of representative clones demonstrated that transcription of $\alpha_2 \delta_1$ mRNA could be reduced to <5% of control in myoblasts and <2% of control in myotubes transduced with siR1. Two myoblast clones, C14 and C21, were chosen for further analysis, and the results of all of the studies done were identical in both. When only one cell line is presented, we present the data from clone C14 as the representative value. Immunohistochemistry (Fig. 2 B) confirms that in wt cells $\alpha_2 \delta_1$ DHPR has a punctate pattern of expression (23,36), which is dictated by the positioning of CaV1.1 (36) and RyR1 (27). In $\alpha_2\delta_1$ DHPR siRNA knockdown myotubes, there is no $\alpha_2 \delta_1$ DHPR detectable (Fig. 2 *B*), but the absence of the $\alpha_2 \delta_1$ DHPR does not affect the targeting of CaV1.1 or RyR1 (data not shown). Cells transduced with the "control" TRPC3 siRNA, to ascertain possible nonspecific consequences on protein expression, have a wt pattern of $\alpha_2 \delta_1$ DHPR expression (data not shown).

Western blot analysis of heavy SR fractions from control and $\alpha_2\delta_1$ DHPR knockdown myotubes probed for RyR1, α_{1S} DHPR subunit, CSQ, FKBP12, SERCA1, and skeletal triadin (β -tubulin was used as a loading control) demonstrates that there is no detectable $\alpha_2\delta_1$ DHPR protein in either of the two clones used in this study but that the absence of the $\alpha_2\delta_1$ subunits was not associated with any differences in the expression of any other triadic protein tested in either myoblasts or myotubes (Fig. 2 *C*). Cells transduced with the "control" TRPC3 siRNA have *wt* amounts of $\alpha_2\delta_1$ DHPR protein as well as RyR1, CaV1.1, CSQ, FKBP12, SERCA1, and skel-

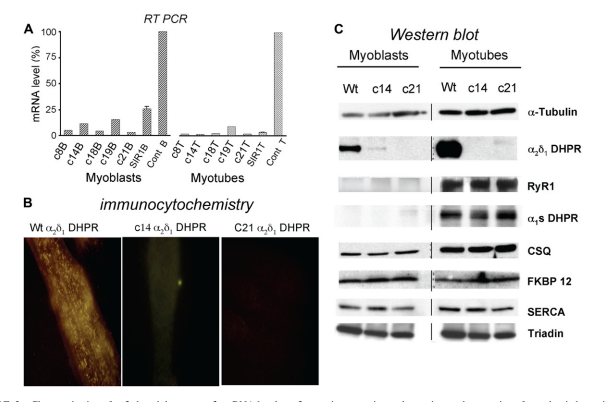


FIGURE 2 Characterization of $\alpha_2\delta_1$ knockdown on $\alpha_2\delta_1$ mRNA levels, $\alpha_2\delta_1$ protein expression and targeting, and expression of muscle triad proteins. (*A*) Transcription of $\alpha_2\delta_1$ mRNA is significantly knocked down in myoblasts and myotubes of both clones that were selected for further study. (*B*) Immunohistochemistry with anti- $\alpha_2\delta_1$ antibody demonstrates that in control myotubes, $\alpha_2\delta_1$ DHPR is targeted appropriately with the characteristic punctate appearance near the cell surface, similar to the staining pattern for α_{1S} DHPR or RyR1 (27). Myotubes from $\alpha_2\delta_1$ knockdown clones 14 and 21 have no visible $\alpha_2\delta_1$ subunit expression. (*C*) Western blot analysis of myoblasts (*left*) and myotubes (*right*) from *wt* and $\alpha_2\delta_1$ knockdown clones 14 and 21. Note that $\alpha_2\delta_1$ is expressed at high levels in *wt* myoblasts and myotubes, whereas α_{1S} DHPR is expressed only in myotubes. The expression of $\alpha_2\delta_1$ protein in knockdown clones 14 and 21 myotubes traces of protein and transcript were essentially not expressed in clone 21 myoblasts or clones 14 and 21 myotubes. The lack of expression of $\alpha_2\delta_1$ appears to have no effect on the expression of any other triadic protein probed.

etal triadin on Western blot analysis (data not shown), showing that the phenotype of the $\alpha_2\delta_1$ knockdown is not simply the result of a nonspecific "off target" artifact caused by siRNA expression.

The effects of $\alpha_2 \delta_1$ knockdown on L-type Ca²⁺ current

To determine the effects of $\alpha_2 \delta_1$ knockdown on the function of the DHPR complex as Ca²⁺ channel, the whole-cell patchclamp configuration was used to measure membrane Ca²⁺ currents (Fig. 3, *A*–*C*). Comparison of representative records demonstrates that, as reported previously (23), currents from $\alpha_2\delta_1$ knockdown myotubes (Fig. 3 *A*) activate more rapidly than those from normal myotubes (Fig. 3 *B*). Activation of currents from normal myotubes was best fitted with the sum of two exponential functions, with a dominant contribution by the slower of these two. The average time constants for these two exponential functions (7.6 ± 0.9 ms and 71.0 ± 3.8 ms; Table 1) were similar to those found previously for normal myotubes (8.5 ± 1.2 ms and 79.5 ± 10.5 ms) by Avila and Dirksen (37). Activation of currents from $\alpha_2\delta_1$ knockdown myotubes could be well fitted with a single exponential function having a time constant (6.4 ± 0.3 ms) similar to that of the faster component for *wt* myotubes (Table 1). At the potential evoking maximal inward current, $\alpha_2\delta_1$

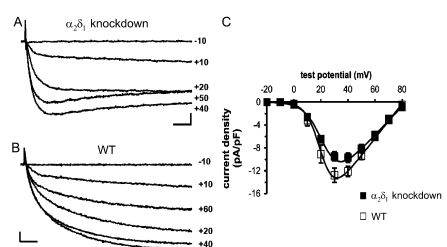
TABLE 1 Characteristics of whole cell Ca²⁺ currents in *wt* and primary $\alpha_2\delta_1$ knockdown myotubes

	$G_{\rm max}~({\rm pS/pF})$	$V_{1/2} ({ m mV})$	$V_{\rm r}~({\rm mV})$	k	Inactivation at 200 ms	$ au_{ m slow}~(m ms)$	$ au_{\mathrm{fast}}~(\mathrm{ms})$	$A_{\rm slow}~({\rm pA/pF})$	A _{fast} (pA/pF)
$Wt \ (n=9)$	307 ± 24	28.0 ± 1.5	81.1 ± 1.3	5.1 ± 0.1	NA	71.0 ± 3.8	7.6 ± 0.9	9.2 ± 0.8	3.7 ± 1.2
$\alpha_2 \delta_1$ k.d. (<i>n</i> = 18)	$248\pm18^*$	22.5 ± 1.3	83.1 ± 1.4	$6.5\pm0.3*$	$(25 \pm 4)\%$	NA	6.4 ± 0.3	NA	9.2 ± 0.8

Current time courses of $\alpha_2 \delta_1$ knockdown cells were fitted with a single exponential function, and current time courses of *wt* myotubes were fitted with a double exponential function.

*p < 0.05 vs. *wt*.

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+30

FIGURE 3 Comparison of Ca²⁺ currents in $\alpha_2\delta_1$ knockdown and normal myotubes. Representative currents elicited by 200 ms test pulses to the indicated potentials are shown from $\alpha_2\delta_1$ knockdown (*A*) and normal (*B*) myotubes. Scale bars are 2 pA/pF (*vertical*) and 20 ms (*horizontal*). (*C*) Average voltage dependence of peak current densities for $\alpha_2\delta_1$ knockdown (closed symbols, n = 18) and normal myotubes (*open symbols*, n = 9). Smooth curves represent the Boltzmann equation (see Materials and Methods) with average parameters (Table 1) obtained from best fits of the same equation to current-voltage relationships of individual cells.

knockdown currents inactivated on average by $25\% \pm 4\%$ at the end of a 200-ms test pulse, whereas no inactivation was observed for *wt* myotubes (Table 1). Peak current-voltage relationships were similar for $\alpha_2\delta_1$ knockdown and normal myotubes, although G_{max} was somewhat lower for $\alpha_2\delta_1$ knockdown myotubes (Fig. 3 *C*; Table 1).

$\alpha_2 \delta_1$ knockdown does not affect DHPR complex tetrad formation and intratetrad spacing

Clusters of DHPR-complex tetrads are equally visible in control and $\alpha_2\delta_1$ DHPR knockdown cells (Fig. 4). Indeed in this particular set of experiments the clusters in the controls (Fig. 4 *A*) tended to contain fewer and less complete tetrads than those in the treated cells (Fig. 4 *B*). Since formation of DHPR-complex tetrads requires an interaction with RyR1, it is clear that $\alpha_2\delta_1$ -null DHPR complexes are appropriately targeted to the calcium release units (CRUs) and appropriately positioned relative to RyRs.

The apparent size of the particles, as indicated by the ring of platinum deposited around them, is not different in the two sets of cells (Fig. 4, A and B). Most importantly, the center-tocenter distance between adjacent particles in the tetrads, or intratetrad distance, is also not altered by the absence of $\alpha_2 \delta_1$. Fig. 4 C shows a scattergram of the measured intratetrad distances, each represented by one dot, blue for controls and orange for treated cells. The x axis of this scattergram is the tetrad number, ranging from 1 to 790. Despite the scatter in the data, due to the distortion that can occur during fracturing, the two average intratetrad distances are the same (19.1 \pm 1.9 nm for control and 19.0 \pm 1.8 nm for $\alpha_2 \delta_1$ DHPR-null cells), and Student's t-test indicates that the two populations are indistinguishable (p = 0.25). This would strongly suggest that 1), the CaV1.1 and not the $\alpha_2 \delta_1$ subunit is the protein which is observed as the large intramembrane particle when the sarcolemma is fractured; and 2), the $\alpha_2\delta_1$ subunit, which is proposed to be a largely extracellular protein, plays no role in the appearance of this particle, in tetrad formation, or in the position of the complexes, as indicated by the intratetrad distance.

$\alpha_2 \delta_1$ is not essential for EC coupling but is important for maintaining Ca²⁺ transients during sustained depolarization

 $\alpha_2 \delta_1$ knockdown cells, in which the mRNA levels were reduced to <2% of control, produced Ca²⁺ transients in response to caffeine which were not significantly different from wt in terms of the peak amplitude (0.75 \pm 0.02 F_{max} for wt and 0.76 \pm 0.3 F_{max} for $\alpha_2 \delta_1$ knockdown), rate of decay (data not shown), and concentration for half-maximal activation (EC₅₀ = 4.6 mM in $\alpha_2 \delta_1$ knockdown vs. 4.6 mM for *wt*). Moreover, $\alpha_2 \delta_1$ knockdown had no significant effect on the amplitude and rate of rise of the Ca²⁺ transient after depolarization with 60 mM KCl (Fig. 5). However, the rate of decay of the Ca²⁺ transient during maintained depolarization was significantly more rapid in $\alpha_2 \delta_1$ knockdown myotubes than in wt. The half-decay time measured as the time (x) axis width of the transient at half-maximum tension fell from 24.7 \pm 1.5 s in *wt* to 5.9 \pm 0.5 s in $\alpha_2\delta_1$ knockdown myotubes (Fig. 5). With this exception the responses from myotubes permanently transduced with the control siRNA were indistinguishable from those seen in controls, demonstrating that the cause of the decrease in the duration of the Ca²⁺ transient and by inference all the other findings were not due to nonspecific "off target" siRNA effects.

Importantly, the difference in the rate of decay of the Ca²⁺ transient during KCl depolarization between $\alpha_2\delta_1$ knockdown and *wt* myotubes could be abolished if extracellular Ca²⁺ was reduced to 7 μ M and extracellular Mg²⁺ increased to 3 mM to maintain the concentration of external divalent cations. Under these conditions, the rate of decay of the Ca²⁺ transient in both groups of myotubes was similar to that of $\alpha_2\delta_1$ knockdown myotubes in Ca²⁺-replete medium (mean $\frac{1}{2}$ decay time of ~5.9 s) (Fig. 6). This strongly suggests that a loss of external Ca²⁺ entry during the 30 s depolarizing

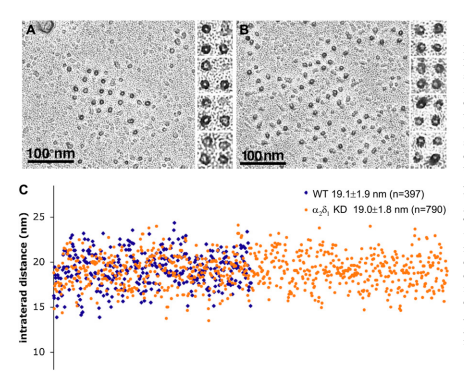


FIGURE 4 Tetrads are unaffected by $\alpha_2\delta_1$ knockdown. (A and B) A dark ringlet of platinum surrounds the freeze fracture particles, each marking the position of the DHPR. Particles and their disposition into tetrads appear identical in the control (*left*) and $\alpha_2 \delta_1$ knockdown cells, indicating that the $\alpha_2 \delta_1$ subunit is not part of the DHPR particle seen using this technique. (C) The center-to-center distance between particles constituting tetrads (intertetrad distance) was measured in electron micrographs at a magnification of \sim 85,000 \times . The tetrads to be measured were selected because they all show a minimum of distortion as a result of fracturing. The images from experimental and control cells were mixed during the analysis so that measurements were performed by a "blind" operator. The data are presented in a scattergram in which each dot represents a single measurement and the x axis is the dot number from 1 to 790. There were no differences in intratetrad distance between the two groups (p > 0.7). All myotubes were fixed in glutaraldehyde, cryoprotected in glycerol, freeze fractured, and rotary shadowed at 45°.

stimulus contributes to the reduced duration of the Ca²⁺ transient in the $\alpha_2\delta_1$ knockdown myotubes.

The relationship between stimulation frequency and Ca²⁺ transient amplitude was studied with pulse trains ranging from 1 to 120 Hz (3 ms duration) and was not found to differ between *wt* and $\alpha_2\delta_1$ knockdown myotubes at any these frequencies (data not shown). Fig. 7 *A* shows that the am-

plitudes of Ca²⁺ transients produced by short electrical stimuli are not significantly different between wt and $\alpha_2\delta_1$ knockdown myotubes. However, unlike wt myotubes, $\alpha_2\delta_1$ knockdown myotubes failed to maintain the peak amplitude of their Ca²⁺ transients for a prolonged, full 120 s, pulse train when the stimulus frequency was increased above 1 Hz. The rate of decay was directly related to the stimulus frequency

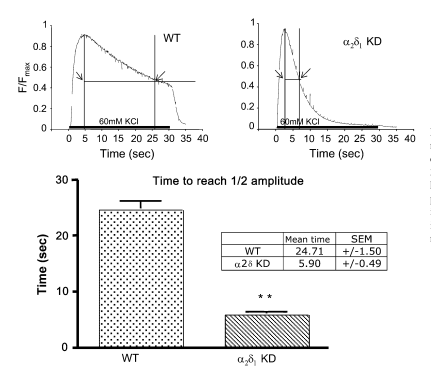


FIGURE 5 Compared to *wt* myotubes, $\alpha_2\delta_1$ knockdown myotubes have abbreviated Ca²⁺ transients during KCl depolarization. Representative, raw Fluo-4 signals during a 30-s 60-mM KCl depolarization of *wt* (*left*) and $\alpha_2\delta_1$ knockdown (*right*) myotubes. The bar graphs in the bottom panel show that the half-decay time for the Fluo-4 transient in response to a 30-s 60-mM KCl depolarization is significantly shorter in $\alpha_2\delta_1$ knockdown myotubes than in *wt* myotubes. Data are shown as mean \pm SE. **p < 0.01; n = 130 myotubes for *wt* and n = 240 for $\alpha_2\delta_1$ knockdown.

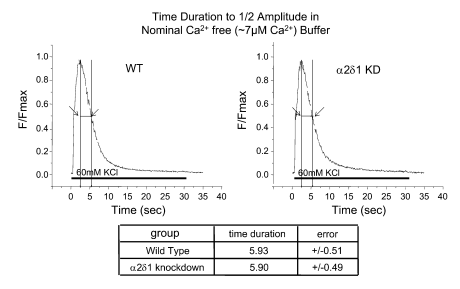
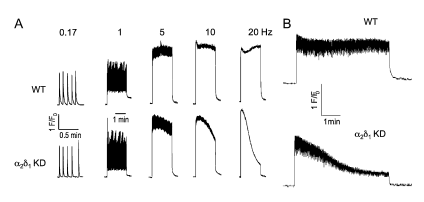


FIGURE 6 $\alpha_2\delta_1$ knockdown and *wt* myotubes have Ca²⁺ transients of similar duration in low Ca²⁺ buffer. Representative average raw Fluo-4 signals during 30-s 60-mM KCl depolarization of n = 65 wt (*left*) and $n = 72 \alpha_2 \delta_1$ knockdown (*right*) myotubes when the depolarization is done in a buffer containing $\sim 7 \ \mu M \ Ca^{2+}$ and 3 mM Mg²⁺. The summary below shows the average time to $\frac{1}{2}$ amplitude \pm mean \pm SE. There is no significant difference between the two genotypes under these conditions. p > 0.8.

(Fig. 7 *A*). Cells were also tested for the rate of fall in their peak Ca²⁺ transient in response to a stimulus protocol similar to those used on adult skeletal muscle cells to produce fatigue. Fig. 7 *B* shows that the amplitude of the Ca²⁺ transient was unaffected during a long (5 min) pulse train composed of bursts lasting 500 ms interspersed by a 500 ms rest period as described in Materials and Methods in *wt* cells. By contrast, $\alpha_2\delta_1$ knockdown cells subjected to the same stimulus protocol had a steady decline in the amplitude of their Ca²⁺ transient. Thus we conclude that the $\alpha_2\delta_1$ DHPR subunit is physiologically necessary for sustaining Ca²⁺ transients in response to prolonged depolarization or repeated trains of action potentials.

$\alpha_2 \delta_1$ knockdown inhibits ECCE

We next tested the hypothesis that the mechanism for the more rapid decay of Ca^{2+} transient amplitude in $\alpha_2\delta_1$ knockdown myotubes was that $\alpha_2\delta_1$ is needed to maintain ECCE, SOCE, or both. To test the effects of $\alpha_2\delta_1$ knockdown on SOCE, we depleted SR Ca^{2+} stores completely with TG while exposing the cells to nominally Ca^{2+} -free external medium. After store depletion was confirmed with the absence of any Ca^{2+} response to exposure to 20 mM caffeine, addition of



2 mM Ca²⁺ to the external medium clearly showed the presence of SOCE (as measured by the Ca2+ transient that accompanied the return to Ca^{2+} -replete medium) (Fig. 8). Using this depletion protocol, the same rate and amplitude of SOCE was measured in both wt and $\alpha_2 \delta_1$ knockdown myotubes (Fig. 9 A, upper traces). In a separate set of experiments, Ca^{2+} depleted myotubes were perfused with an external medium containing 40 mM K⁺ and 2 mM Ca²⁺ (Fig. 9 A, lower *traces*). K^+ depolarization elicited rapid Ca²⁺ entry in *wt* cells via ECCE, whereas $\alpha_2 \delta_1$ knockdown myotubes failed to show any detectable Ca²⁺ entry while the cells were depolarized. When K^+ was removed from the Ca^{2+} -replete external medium, a rapid Ca²⁺ entry was observed in the $\alpha_2 \delta_1$ knockdown myotubes similar to their previous response with no depolarization (Fig. 9 A, lower traces). Thus the abolition of ECCE during KCl depolarization in the $\alpha_2 \delta_1$ knockdown myotubes allowed the direct demonstration that SOCE is completely inhibited by sarcolemmal depolarization.

We also examined Mn^{2+} quench of fura-2 fluorescence during electrical (20 Hz trains) and 40 mM K⁺ depolarization. Fig. 9 *B* shows that the rate of Mn^{2+} entry is significantly reduced (p < 0.0003) in response to either KCl or electrical depolarization in $\alpha_2\delta_1$ knockdown myotubes.

> FIGURE 7 $\alpha_2 \delta_1$ knockdown myotubes fail to maintain their Ca²⁺ transient amplitude with electrical pulse trains >1 Hz. (A) Cells were stimulated by sequential bipolar electrical pulse trains lasting 120 s at frequencies ranging from 0.17 to 20 Hz with a 1 min rest period between each frequency tested. The rate of decline of the peak Ca²⁺ transient amplitude was proportional to the stimulus frequency. These data are representative of measurements acquired from n = 3 wt and $n = 8 \alpha_2 \delta_1$ knockdown myotubes using the indicated stimulus sequence protocol. (B) Ca^{2+} transient amplitudes were resistant to rundown in wt, but not $\alpha_2 \delta_1$ knockdown, myotubes stimulated with electrical pulse trains similar to those commonly used to elicit fatigue in adult fibers, as described in Materials and Methods. The data are representative of n = 20 wt and n =36 knockdown cells.

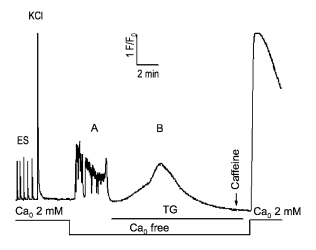


FIGURE 8 Complete SR Ca depletion protocol for $\alpha_2\delta_1$ knockdown myotubes. Cells were loaded with Fluo-4 as described in Materials and Methods and tested for responses to electrical pulses and K⁺ (40 mM) application in Ca-replete medium (Ca₀²⁺ 2 mM). The extracellular solution was exchanged with one lacking added Ca²⁺ (Ca₀²⁺ free); it elicited spontaneous activity in most cells. TG (200 nM) was then perfused onto the cells in the Ca₀²⁺-free medium until the new baseline was established (typically <10 min) and depletion of SR Ca tested with a 20 mM caffeine challenge. The presence of SOCE was then tested by perfusion of 2 mM Ca²⁺ medium. This protocol completely depleted SR Ca in >95% of all myotubes exhibiting EC coupling. *Wt* myotubes were indistinguishable from the $\alpha_2\delta_1$ knockdown myotubes shown here.

The rate of Mn^{2+} entry is reduced by more than twofold in response to electrical stimulation, and it is abolished in response to K^+ depolarization.

ECCE requires the presence of RyR1 and CaV1.1 and is very dependent on the conformation of RyR1. We have shown that ECCE can be enhanced by two conformational mutations of RyR1 that do not support normal EC coupling (4) or by pretreatment of murine myotubes (4,5) or muscle fibers (12) with 250–500 μ M ryanodine. This apparent enhancement of ECCE elicited by changes in the conformation of RyR1 caused by mutations or ryanodine binding is primarily a result of delayed inactivation/deactivation of entry during depolarization and not increased activation. To test the hypothesis that the $\alpha_2 \delta_1$ subunit was key to the interaction of the three Ca²⁺ channels participating in ECCE, we pretreated $\alpha_2 \delta_1$ knockdown myotubes with either 500 μ M ryanodine or solvent (control) for 30 min and performed Mn²⁺ quench experiments similar to those described above. In Fig. 9, C and D, it can be seen that, unlike wt myotubes, 40 mM K^+ depolarization of $\alpha_2 \delta_1$ knockdown myotubes in the absence of ryanodine pretreatment failed to permit Mn²⁺ entry. However $\alpha_2 \delta_1$ knockdown myotubes pretreated with ryanodine regained a robust long-lived K⁺-triggered Mn²⁺ entry response, although the initial rate was much ($\sim 60\%$) smaller than the rate seen in wt (Fig. 9 D). This would indicate that although the $\alpha_2 \delta_1$ subunit is required for physiologic ECCE, its absence can be partially overcome when RyR1 is put into a conformation that enhances this depolarization-induced Ca^{2+} entry.

DISCUSSION

$\alpha_2 \delta_1$ is not necessary for myotube growth or differentiation

Despite the almost complete absence (<2%) of the $\alpha_2\delta_1$ DHPR subunit, there appears to be no change in the ability of the myoblasts to form myotubes or in the expression levels or targeting of CaV1.1 to the CRU (28,38). This is similar to the report of Obermair et al. (23), who studied transiently siRNA transfected cells with a reduction in $\alpha_2\delta_1$ mRNA to 6%–25% of control. In addition, the absence of the $\alpha_2\delta_1$ DHPR subunit did not affect the expression of several other components of the triad (RyR1, FKBP12, CaV1.1, CSQ, Triadin, SERCA), suggesting that the $\alpha_2\delta_1$ subunit is not important for the expression or normal targeting of these CRU proteins.

Unlike CaV1.1, the $\alpha_2\delta_1$ subunit is expressed in both myoblasts and myotubes, and $\alpha_2\delta_1$ -null mice have an embryonic or birth lethal phenotype (J. Offord, Pfizer, Groton, CT, personal communication, 2007), suggesting that the $\alpha_2\delta_1$ may play a role in embryonic development of components other than muscle. Our data show that whatever the early function of $\alpha_2\delta_1$ in myoblasts, it is not necessary for myoblast division, their differentiation into myotubes, or basic EC coupling.

The effects of the absence of the $\alpha_2 \delta_1$ subunit on slow voltage-gated Ca^{2+} current

Overall, our results on Ca²⁺ currents in $\alpha_2\delta_1$ knockdown myotubes are in good agreement with those reported previously showing that the fundamental functions of the DHPR complex as Ca²⁺ channel do not depend on the presence of the $\alpha_2\delta_1$ subunit (23). In particular, $\alpha_2\delta_1$ knockdown had little effect on the amplitude or voltage dependence of L-type Ca²⁺ currents but did result in accelerated activation kinetics. Additionally, in response to 200-ms depolarizations, the L-type currents displayed greater inactivation in $\alpha_2\delta_1$ knockdown myotubes, perhaps because inactivation in *wt* myotubes was obscured by the slower activation kinetics.

Lack of the $\alpha_2 \delta_1$ subunit reveals the position of CaV1.1 in DHPR complex tetrads

Freeze fracture reveals the position of intramembranous proteins, but it cannot be reliably used to define the size and shape of the proteins that are fractured in the frozen state. For example, the very large ryanodine receptor of the SR produces a barely visible shallow hump when fractured (39). Nevertheless, we are surprised to find that even though the $\alpha_2\delta_1$ is a very large subunit, the DHPR complex particles are neither absent nor grossly distorted in its absence. The importance of this result lies in the fact that we can, albeit at a low level of resolution, predict the position of the α_{1s} DHPR mass relative to the four RyR subunits (40). The channel would be located somewhat peripherally relative to the RyR

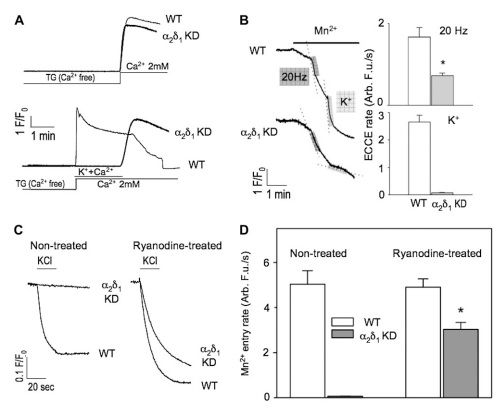


FIGURE 9 $\alpha_2 \delta_1$ knockdown selectively inhibits ECCE but not SOCE. (A) (Upper traces) $\alpha_2\delta_1$ knockdown had no effect on initial rate and amplitude of SOCE under condition of severe store depletion elicited by TG (200 nM). (Lower traces) ECCE is completely abolished in K⁺-depolarized $\alpha_2 \delta_1$ knockdown myotubes. Note that K⁺ depolarization in the presence of external Ca²⁺ reveals the complete inhibition of SOCE even under conditions of severe depletion. (B) Representative traces showing Mn2+ quench of fura-2 fluorescence in $\alpha_2 \delta_1$ knockdown cells was significantly reduced when a 20 Hz electrical pulse train was applied or completely abolished when the cells were exposed to 60 mM KCl. (C) Representative traces of KCl depolarization induced Mn2+ influx pre- and posttreatment with ryanodine. Although there is a brisk Mn^{2+} influx in *wt* cells when they are exposed to 60 mM KCl, there is no Mn^{2+} influx in $\alpha_2\delta_1$ knockdown cells under normal conditions. However, when the cells are pretreated with high concentrations of ryanodine (500 μ M) for 30 min ECCE can be partially rescued ($\sim 40\%$ of wt) by the change in

the conformation of RyR1 caused by ryanodine exposure. (D) Even in the presence of ryanodine, the amount of Mn^{2+} entry in $\alpha_2\delta_1$ knockdown cells is significantly reduced compared to *wt*. N > 20 for each group.

outline in such a manner that some portion of it could extend over the side of RyR1.

$\alpha_2 \delta_1$ is not essential for EC coupling, but its absence alters properties of the Ca²⁺ transient

In agreement with Obermair et al. (1), we find that the $\alpha_2 \delta_1$ DHPR subunit is not needed for skeletal-type EC coupling since there was no difference between $\alpha_2 \delta_1$ -deficient and wt myotubes in either the magnitude or the activation kinetics of the Ca²⁺ transient elicited by electrical or KCl depolarization in nominally Ca^{2+} -free buffer (7 μ M). However, during sustained depolarization in Ca^{2+} -containing (2 mM) medium, the duration of Ca^{2+} transients in $\alpha_2\delta_1$ knockdown myotubes was reduced \sim 5-fold compared to *wt* myotubes, despite similar sized SR Ca²⁺ stores, as indicated by caffeine responses. Two observations support the idea that this reduced duration in $\alpha_2 \delta_1$ knockdown cells is a consequence of a loss of Ca^{2+} entry via ECCE. First, measurements of Mn^{2+} quench indicate that ECCE is greatly reduced in $\alpha_2 \delta_1$ knockdown myotubes. Second, the duration of Ca²⁺ transients in wt myotubes is decreased fivefold when the external medium is made nominally Ca²⁺-free (Mg²⁺ increased to 3 mM), whereas this manipulation has no effect on the duration of the Ca^{2+} transient in $\alpha_2\delta_1$ knockdown cells. Although these results point to the importance of Ca^{2+} entry via ECCE

for maintaining Ca²⁺ transients during prolonged depolarization, two limitations should be mentioned. First, removal of external Ca²⁺ would eliminate all sources of Ca²⁺ entry (not just that via ECCE). Second, removal of external Ca²⁺ might be expected to cause the voltage sensor for EC coupling to inactivate more rapidly (41,42). However, if either of these represented major mechanisms whereby Ca²⁺ removal caused a reduced duration of the Ca²⁺ transient in *wt* myotubes, it would not explain why Ca²⁺ removal had essentially no effect on $\alpha_2\delta_1$ knockdown cells.

$\alpha_2 \delta_1$ is essential for physiologically engaging ECCE but not SOCE

In cells treated with TG, there was no difference in the kinetics or magnitude of the refilling response between $\alpha_2\delta_1$ DHPR knockdown and *wt* myotubes, showing that the absence of the $\alpha_2\delta_1$ subunit has no detectable effect on SOCE. Interestingly, when another set of myotubes from the same myoblast clones were depolarized with 40 mM KCl concurrently with the addition of 2 mM Ca²⁺ in the external medium, there was a striking difference between the two cell types. As expected (5), *wt* cells had a rapid Ca²⁺ transient with an even faster rise time than was seen with the addition of Ca²⁺ alone (5,43–45). $\alpha_2\delta_1$ -deficient cells, on the other hand, showed no Ca²⁺ entry response at all during the KCl

depolarization but did show a rate of cation entry similar to the rate of the SOCE transient when the KCl was removed. This result suggests that the mechanism behind the rapid decline in the Ca²⁺ transient during a 30 s KCl exposure or trains of 20 Hz electrical stimuli in $\alpha_2\delta_1$ knockdown myotubes was either a marked reduction or a complete absence of ECCE. In addition these data directly demonstrate and confirm the suggestions by others that SOCE is significantly inhibited by depolarization (46–48).

The results of our studies using Mn²⁺ quench of fura-2 fluorescence clearly confirm that that $\alpha_2 \delta_1$ is an essential regulator of physiologic ECCE in skeletal myotubes and the degree of the dysfunction depends on 1), the nature of the depolarizing trigger (electrical trains versus [K⁺]), and, importantly, 2), the conformation of RyR1. From our results, it appears that if the RyR1 Ca²⁺ release channel is in or is placed into a conformational state different from the state found at rest in the cell (e.g., with the conformational state induced by certain RyR1 mutations or ryanodine), the effect of the absence of $\alpha_2 \delta_1$ on ECCE can be partially overcome. This observation makes it unlikely that the absence of $\alpha_2 \delta_1$ has either negatively influenced the expression or targeting of the ECCE channel to the surface membrane. Presumably the reason that ECCE is not completely abolished in $\alpha_2 \delta_1$ -deficient myotubes during electrical stimulation is that at a fairly low frequency of stimulation the voltage sensor has the opportunity to reset itself between individual stimuli. It is possible that if the stimulus frequency were raised higher than 20 Hz the reduction in the rate of Mn^{2+} entry would be more profound. This would suggest that the physiological engagement of ECCE may be most significant under conditions of extreme activity and that changes in the efficiency of Ca²⁺ entry during depolarization may be a factor in the processes leading to muscle fatigue.

In conclusion, the evidence gained from these studies confirms the previously defined role of the $\alpha_2 \delta_1$ DHPR subunit as the particle needed to stabilize the CaV1.1 channel in the slowly activating state. Further it provides conclusive evidence that the $\alpha_2 \delta_1$ subunit is not the particle that forms tetrads in freeze fracture studies of DHPRs in the surface membrane. Most importantly, it demonstrates that in addition to its previously defined role, the $\alpha_2 \delta_1$ subunit also plays an important role in the three-way interaction of the DHPR complex, RyR1, and ECCE channels during normal EC coupling. However, unlike the CaV1.1, the role of the $\alpha_2\delta_1$ subunit in coordinating ECCE is important but not essential, as the absence of $\alpha_2 \delta_1$ can be partially overcome by repriming the voltage sensor and almost completely overcome when the RyR1 assumes a conformation that enhances ECCE.

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