Muscle-specific RING finger-1 interacts with titin to regulate sarcomeric M-line and thick filament structure and may have nuclear functions via its interaction with glucocorticoid modulatory element binding protein-1

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he COOH-terminal A168–170 region of the giant sarcomeric protein titin interacts with muscle-specific RING finger-1 (MURF-1). To investigate the functional significance of this interaction, we expressed green fluorescent protein fusion constructs encoding defined fragments of titin's M-line region and MURF-1 in cardiac myocytes. Upon expression of MURF-1 or its central region (containing its titin-binding site), the integrity of titin's M-line region was dramatically disrupted. Disruption of titin's M-line region also resulted in a perturbation of thick filament components, but, surprisingly, not of the NH2-terminal or I-band regions of titin, the Z-lines, or the thin filaments. This specific phenotype also was caused by the expression of titin A168–170. These data suggest that the interaction of titin with MURF-1 is important for the stability of the sarcomeric M-line region.

MURF-1 also binds to ubiquitin-conjugating enzyme-9 and isopeptidase T-3, enzymes involved in small ubiquitin-related modifier–mediated nuclear import, and with glucocorticoid modulatory element binding protein-1 (GMEB-1), a transcriptional regulator. Consistent with our in vitro binding data implicating MURF-1 with nuclear functions, endogenous MURF-1 also was detected in the nuclei of some myocytes. The dual interactions of MURF-1 with titin and GMEB-1 may link myofibril signaling pathways (perhaps including titin's kinase domain) with muscle gene expression.

Introduction

Numerous structural and regulatory proteins are assembled and maintained in an exquisitely precise order within the myofibrils of striated muscle, yet the molecular mechanisms responsible for this phenomenon are not well understood. The principal components of muscle sarcomeres, the basic contractile units of myofibrils, include parallel arrays of actin-containing thin filaments that overlap with myosin-containing thick filaments. A third filament system is formed by single molecules of titin (connectin), the largest vertebrate protein identified to date (mol wt ~3–3.7 MD) (Maruyama et al., 1977; Wang et al., 1979; Labeit and Kolmerer, 1995; Bang et al., 2001). Titin molecules from adjacent sarcomeres overlap in the Z-line, and those from opposite half sarcomeres overlap in the M-line, thus forming a continuous filament system within myofibrils (Obermann et al., 1996; Gregorio et al., 1999). The majority (~90%) of the titin molecule is comprised of repeating modular domains from the fibronectin (FN)* type III and the immunoglobulin (Ig) superfamilies. Additionally, there are 17 unique

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*Abbreviations used in this paper: FN, fibronectin; GFP, green fluorescent protein; GMEB-1, glucocorticoid modulatory element binding protein-1; GST, glutathione-S-transferase; Ig, immunoglobulin; ISOT-3, isopeptidase T-3; MURF-1, muscle-specific RING finger-1; MyBP-C, myosin-binding protein C; RBCC, RING finger-B-box-coiled-coil; RNF, RING finger; SMRZ, striated muscle RING zinc finger; SUMO, small ubiquitin-related modifier; Ubc9, ubiquitin-conjugating enzyme 9; Y2H, yeast two-hybrid.
insertions distributed along the length of titin, including a serine/threonine kinase domain in its M-line region (Labeit et al., 1992), the function of which remains elusive.

To dissect the functional properties of titin’s domains, recent investigations have focused on novel titin-binding proteins. One of these proteins is muscle-specific RING finger-1 (MURF-1) (Centner et al., 2001), also recently identified as striated muscle RING zinc finger (SMRZ) (Dai and Liew, 2001) and RING finger 28 (RNF 28) (see http://www.gene.ucl.ac.uk/cgi-bin/nomenclature/searchgenes.pl). This protein binds to the titin domains A168–169–170 (A168–170) located directly NH₂-terminal to the titin kinase domain (Centner et al., 2001). Two proteins with a high degree of homology to MURF-1 also have been identified: MURF-2 (RNF 29) and MURF-3 (RNF 30) (Spencer et al., 2000; Centner et al., 2001). The three MURFs are members of the RING finger-B-box-coiled-coil (RBCC) family, a class of proteins that have critical roles in cellular processes including signal transduction, gene transcription, ubiquitination, and differentiation (for reviews see Freemont, 2000; Borden, 2000). Structurally, the MURFs contain a Zn-binding RING finger domain at their extreme NH₂-terminal end, a MURF family–specific conserved region, a B-box domain, coiled-coil motifs, and an acidic tail (Spencer et al., 2000; Centner et al., 2001; Dai and Liew, 2001).

To date, little insight into the cellular roles of the MURFs is available. In vitro binding studies revealed that MURF family members homo- and hetero-oligomerize (Centner et al., 2001). MURF-3 appears to associate with microtubules and have a role in myogenic differentiation and microtubule stabilization (Spencer et al., 2000). MURF-1 (SMRZ) recently has been shown to interact with small ubiquitin-related modifier-3 (SUMO-3/SMT3b; Dai and Liew, 2001), a member of a ubiquitin-related class of proteins implicated in subcellular targeting and nuclear import (for review on SUMO proteins see Melchior, 2000). Consistent with this finding, MURF-1 is detected in nuclei (Dai and Liew, 2001; this study). In a different study, it was determined that MURF-1 appears to be the only MURF family member that interacts directly with titin, within the M-line region of the sarcomere (Centner et al., 2001). Despite these recent studies, the exact physiological role(s) of the MURF family members, particularly MURF-1 and -2, have remained elusive.

To determine which regions of MURF-1 target to myofibrils and/or nuclear sites, and as an initial approach to decipher the cellular properties of MURF-1, we expressed green fluorescent protein (GFP) fusion constructs encoding defined regions of MURF-1 and its titin-binding site, A168–170, in live cardiac myocytes. Our data suggest that the interaction of titin with the central region of MURF-1 is important for maintaining the integrity of titin’s M-line structure. In turn, titin’s COOH-terminal (M-line) region appears to be necessary for thick filament integrity, but, surprisingly, not for the integrity of titin’s NH₂-terminal or I-band region, the thin filaments, or the Z-lines.

Intriguingly, endogenous MURF-1 also was detected in the nuclei of some myocytes. Consistent with this observation, in vitro interaction studies revealed that MURF-1 binds to ubiquitin-conjugating enzyme 9 (Ubc9) and isopeptidase T-3 (ISOT-3), enzymes involved in SUMO modification of target proteins, a posttranslational modification that occurs in the nucleus. Our in vitro interaction studies also demonstrated that MURF-1 is capable of binding to glucocorticoid modulatory element binding protein-1 (GMEB-1), a nuclear protein implicated in transcriptional regulation (Theriault et al., 1999; Jimenez-Lara et al., 2000; Zeng et al., 2000b). Therefore, our data suggest MURF-1 has an important role in titin filament M-line structure (and perhaps in titin kinase-based signaling processes) as well as a nuclear function (potentially in the control of muscle gene expression). Future studies will likely provide insights into how the dual functions of MURF-1 are associated, as well as how myofibrillogenesis and the regulation of muscle gene expression are linked.

Results

MURF-1 has multiple subcellular localizations in cardiac myocytes.

Previous immunolocalization studies revealed that in adult striated muscle, MURF-1 is distributed diffusely throughout the cytoplasm, is assembled at the Z-line region, and is at the M-line region where its binding partner, titin A168–170, is located (Centner et al., 2001). We used primary cultures of fetal rat and embryonic chick cardiac myocytes in studies de-

![Figure 1](image-url)
signed to investigate the cellular role of the interaction of MURF-1 with titin. When these cells were stained with anti–MURF-1 antibodies, MURF-1 was detected diffusely throughout the cytoplasm, and was assembled at the M-line region (Fig. 1; data from rat myocytes shown). Costaining for α-actinin, which stains all myofibrils, revealed that in some myocytes, MURF-1 assembled in only a few myofibrils, whereas in other myocytes, it was assembled in all myofibrils (Fig. 1 a, arrows). Consistent with this observation, immunolocalization studies on isolated rat cardiac and skeletal muscle myofibrils also demonstrated that MURF-1 was detected at the M-line region in only a portion of myofibrils (30–50%; unpublished data). Interestingly, although MURF-1 was detected at the M- and the Z-line in adult cardiac tissue sections (Centner et al., 2001), MURF-1 was not detected at the Z-line in rat fetal or chick embryonic cardiac myocytes. The significance of this is not known, but may be due to developmental differences. MURF-1 staining was also observed diffuse in the cytoplasm (Fig. 1, c and e, arrowheads) and at varying intensities in the nuclei of some cardiac myocytes, as shown by colocalization with DAPI staining (Fig. 1, e and f); this was confirmed by scanning at different focal planes by deconvolution microscopy (unpublished data). MURF-1's nuclear localization is consistent with a recent study that detected MURF-1 (SMRZ) fusion proteins in the nuclei of C2C12 skeletal muscle cells (Dai and Liew, 2001). Staining with the secondary antibody alone yielded negligible background in all experiments (unpublished data).

Expression of MURF-1 or titin A168–170 disrupts the integrity of titin’s M-line region and the organization of thick filament components.

We generated a GFP–MURF-1 fusion construct for expression studies in chick cardiac myocytes (Fig. 2). Confirming the immunolocalization studies, GFP–MURF-1 assembled in a
portion of myofibrils at the M-line region and colocalized with anti-titin A168–170 staining (unpublished data). GFP–MURF-1 was also diffusely distributed and in distinct aggregates (of varying size and intensity) in the cytoplasm of transfected myocytes (Fig. 3 c, arrowheads), as well as in contaminating fibroblasts that normally do not express MURF-1 (unpublished data). These aggregates are likely a result of homo-oligomerization, which has been reported previously with MURF proteins (Spencer et al., 2000; Centner et al., 2001). Titin staining also colocalized with some GFP–MURF-1 aggregates in the cytoplasm of myocytes (Fig. 3 c and d, arrowheads) perhaps because the MURF-1 aggregates provided additional binding sites for titin during myofibril assembly or turnover. GFP–MURF-1 also was observed in the nuclei of some myocytes (unpublished data). Strikingly, the majority of the GFP–MURF-1–transfected cells exhibited severe disruption of the COOH-terminal region of titin compared with cells expressing GFP alone (Fig. 3, b and d, arrows).

Surprisingly, triple-labeling studies revealed that although titin’s COOH-terminal region was disrupted in GFP–MURF-1–expressing myocytes, titin’s NH₂-terminal region appeared in a regular, striated pattern in the identical myofibrils (Fig. 3, i and j). However, staining transfected cells for myosin, myosin-binding protein C (MyBP-C), and myomesin demonstrated that GFP–MURF-1 expression also perturbed thick filament integrity in a large percentage of transfected myocytes (Fig. 4). Specifically, ~80–90% of cells transfected with MURF-1 had disrupted COOH-terminal titin structure, whereas ~60–70% of the cells had perturbed thick filament structure (Fig. 6). This difference in percentages of cells affected suggests that the thick filament perturbation may be a secondary effect of the disruption of titin’s M-line region. Unexpectedly, the structure of the actin (thin) filaments, the I-band region of titin, and the Z-lines (as assessed by the distribution of phalloidin, titin N2A, and α-actinin staining, respectively) appeared in typi-

Table 1. Subcellular localization of GFP–MURF-1 deletion constructs in cardiac myocytes

<table>
<thead>
<tr>
<th>GFP–MURF-1 deletion construct</th>
<th>Cytoplasmic aggregates</th>
<th>Striated at M-line region</th>
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<tr>
<td>GFP alone</td>
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<td>RING</td>
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<td>Tail</td>
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<td>≤10</td>
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<tr>
<td>Tailless</td>
<td>90–95</td>
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<td>RINGless</td>
<td>90–95</td>
<td>30–40</td>
<td>0</td>
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<tr>
<td>Central</td>
<td>90–95</td>
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*a*Represents the total percentage of transfected myocytes observed to contain GFP fusion proteins in the specified localization.

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**Figure 4.** Expression of GFP–MURF-1 also perturbs the organization of thick filament components. Transfected myocytes were stained with antibodies to myosin (b and d), MyBP-C (f and h), and myomesin (j and l). In many GFP–MURF-1–expressing myocytes, staining for thick filament components was perturbed (d, h, and l), compared with myocytes transfected with GFP alone (b, f, and j). Single arrows mark perturbed thick filament component staining, and double arrows mark regular, striated thick filament staining. Note, size and intensity of the GFP–MURF-1 cytoplasmic aggregates vary from cell to cell (c, g, and k, arrowheads). Bar, 10 μm.
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cal striated patterns in the majority of GFP–MURF-1–transfected cells (Figs. 5 and 6), even though the COOH-terminal region of titin was disrupted (Fig. 5, i and j).

Because MURF-1 was shown to interact with the COOH-terminal titin domains A168–A170 (Centner et al., 2001), we also performed expression studies with titin GFP fusion constructs encoding the A168–170 region or its flanking regions for comparison (Fig. 2). When expressing titin A168–170–GFP in myocytes, both the M-line region of titin (Fig. 7 b) and the thick filaments (Fig. 7 h; data shown for myomesin, similar results were obtained for myosin and MyBP-C) were perturbed. Consistent with the results from the GFP–MURF-1 expression studies, Z-line and thin filament structure appeared in their characteristic patterns in the titin A168–170–GFP–transfected cells (Fig. 7, n and t).

In contrast, myocytes expressing other titin M-line regions (the Ig domains located COOH-terminal to the A168–170 region, M1-M2-M3–GFP or M8-M9-M10–GFP) had regular, striated staining patterns for titin, thick filament, Z-line, and thin filament components (Fig. 7, d, j, p, and v; data shown for M8-M9-M10-GFP). Furthermore, myocytes expressing GFP fusion constructs encoding titin’s unique Ser/Thr kinase domain, as well as those expressing a constitutively activated mutant form of this domain, exhibited typical titin, thick filament, thin filament, and Z-line staining (Fig. 7, f, l, r, and x; data shown for titin kinase–GFP).

In conclusion, the disruption of sarcomeric M-line integrity was specific to the expression of titin A168–170 and its ligand, MURF-1. These data suggest that the interaction of titin with MURF-1 has an important role in maintaining the structure of sarcomeric M-line components in cardiac myocytes. Our data also surprisingly indicate that disruption of the integrity of the COOH-terminal region of titin perturbs thick filament structure but does not appear to affect the structure of titin’s I-band or NH2-terminal regions, the thin filaments, or the Z-lines.

MURF-1’s central region targets to the M-line and maintains M-line structure, whereas the RING domain targets to nuclei

To investigate the regions of MURF-1 involved in its subcellular targeting and maintaining the structure of titin’s COOH-terminal region, we generated five GFP–MURF-1 deletion constructs (Fig. 2). One construct encoded the MURF-1 NH2-terminal RING domain alone (RING), which contains the SUMO-3/SMT3b binding site (Dai and Liew, 2001). Another construct encoded only the COOH-terminal acidic tail region (Tail). Three constructs contained the central region of MURF-1, which binds to titin A168–170 (Centner et al., 2001): one encoded the NH2-terminal RING domain plus the central region (Tailless); one encoded the central region

Figure 5. Expression of GFP–MURF-1 does not appear to affect the integrity of thin filament or Z-line components. Myocytes expressing GFP–MURF-1 (c) or GFP alone (a) were stained for Z-lines with sarcomeric α-actinin antibodies (b and d) and show regular, striated staining. Triple-labeling studies in GFP–MURF-1–transfected cells (h), using Texas red–conjugated phalloidin (j) and antibodies to titin A168–170 (i), determined that thin filament integrity is not affected upon disruption of COOH-terminal titin in identical myofibrils. GFP–transfected cells (e) exhibited normal actin filament (g) and COOH-terminal titin (f) staining. Double arrows mark regular, striated staining. Single arrows mark disrupted titin staining. Bars, 10 μm.

Figure 6. Quantification of GFP–MURF-1– and GFP-expressing myocytes exhibiting disrupted staining patterns for sarcomeric components. Myocytes expressing GFP–MURF-1 (gray bars) or GFP alone (white bars) were stained for various sarcomeric components, and the number of cells exhibiting perturbed staining patterns was counted. The results indicate that the COOH-terminal region of titin (A168–170 and AB5) was severely disrupted in most GFP–MURF-1–expressing cells, whereas its I-band (N2A) and NH2-terminal region (T11) were not affected. The thick filament components myosin, myomesin, and C-protein were also disrupted in a large majority of GFP–MURF-1–expressing cells, but thin filament and Z-line components appear relatively unaffected. Data are presented as the mean percentage of total myocytes with disrupted staining ± SD. Means were obtained by counting >50 myocytes from more than two experiments and the results are representative of >10 experiments.
plus the COOH-terminal tail (RINGless); and one encoded the MURF-1 central region alone (Central). Neither RING nor Tail were observed to assemble at the sarcomeric M-line region or form cytoplasmic aggregates. Strikingly, however, RING–GFP was detected in the nuclei of all transfected myocytes observed (Table I). Because aberrant nuclear localization of GFP fusion proteins is known to occur, we quantified the nuclear localization of each GFP–MURF-1 fusion protein in transfected myocytes (Table I). RING–GFP localized to nuclei in 100% of observed cells, compared with GFP alone, which was observed in nuclei in 40–65% of cells. Consistent with this, RINGless–GFP never exhibited nuclear localization, but Tailless–GFP (containing the RING domain) was detected in nuclei in 40–65% of cells (Table I). Finally, Tail–GFP was detected in ≤10% of myocyte nuclei, suggesting that it does not contain a fully functional nuclear targeting site. These data, together with the observations that MURF-1’s RING domain binds to the nuclear protein SUMO-3 (Dai and Liew, 2001) and other RING proteins have been reported in nuclei (for review see Borden, 2000), are consistent with the RING domain targeting MURF-1 to the nuclei of cardiac myocytes.

In contrast, all three constructs encoding MURF-1’s central region (Tailless, RINGless, and Central) assembled at the M-line region in a portion of myofibrils of some myocytes (Table I). Additionally, the expressed proteins all formed cytoplasmic aggregates in the majority of transfected cells, most likely due to their coiled-coil domains. These data indicate that MURF-1’s central region, containing the titin-binding site, targets the protein to the sarcomeric M-line region, whereas the RING domain appears to target it to nuclei.

Next, we costained transfected myocytes with anti–COOH-terminal titin antibodies to identify the region(s) of MURF-1 involved in M-line structure. Expression of each of the three MURF-1 fragments containing its central region, shown above to contain an M-line targeting site, resulted in a phenotype identical to that observed when full-length MURF-1 was expressed. Specifically, a marked disruption of titin A168–170 staining was observed in the vast majority of cells expressing Central, Tailless, or RINGless (Fig. 8, f, h, and j), compared with regular, striated titin staining in myocytes expressing RING or Tail (Fig. 8, b and d). These studies reveal that MURF-1 contains two distinct targeting and functional domains. Its central region, containing the titin-binding site, targets it to the M-line and participates in maintain-
ing the integrity of titin’s COOH-terminal region. The RING domain, containing the binding site for SUMO-3, appears to be involved in MURF-1 nuclear targeting.

**MURF family members interact with Ubc9 and ISOT-3, but only MURF-1 interacts with the transcriptional modulator GMEB-1**

Previous binding studies revealed that MURF-1 interacts with titin A168–A170 and with itself (by oligomerization), and also hetero-oligomerizes with MURF-2 and -3 (Centner et al., 2001). Other in vitro interaction studies determined that MURF-1 also binds to SUMO-3/SMT3b (Dai and Liew, 2001). We searched for additional MURF binding proteins in human heart and fetal mouse cDNA libraries by yeast two-hybrid (Y2H) screening. These novel screens demonstrated that both MURF-1 and -2 baits interact with Ubc9 and ISOT-3 prey clones (Fig. 9 A). Ubc9 is an enzyme involved in SUMO modification; it specifically catalyzes the formation of an isopeptide bond between SUMO

Figure 9. MURF family members interact with SUMO modifying enzymes ISOT-3 and Ubc9, but only MURF-1 interacts with the transcriptional regulator GMEB-1. (A) Y2H screens using full-length cDNAs of individual MURF family members as baits identified ISOT-3 (light gray) and Ubc9 (black) as MURF-binding proteins. However, GMEB-1 (dark gray) was found to interact only with MURF-1. β-Galactosidase assays were performed to confirm positive clones, and the levels were compared with colonies transformed with each prey construct and the empty bait vector (white). Data are presented as mean levels of β-galactosidase from triplicate experiments ± SD. ***, P > 0.001. (B) RT-PCR analysis of human heart total RNA revealed that GMEB-1 mRNA transcripts are detectable in heart (H) and skeletal (Sk) tissues. Lane 1, no reverse transcriptase control in human heart RNA (−); lane 2, 511-bp GMEB-1 PCR product amplified from human heart RNA (+); lane 3, 511-bp GMEB-1 PCR product amplified from human skeletal RNA (−); lane 4, no reverse transcriptase control in human skeletal RNA (−). (C) GMEB-1 specifically binds to MURF-1 in GST pull-down assays. GMEB-1 was translated in vitro (lane 3). When incubated with bacterially expressed GST–MURF-1 fusion peptides, GMEB-1 and MURF-1 binding to glutathione–sepharose 4B beads was detectable (lane 2). Lane 1 contains no detectable binding of GMEB-1 to the beads alone. IVT, in vitro translated. (D) GMEB-1–GFP targets to the nuclei of cardiac myocytes (a) and MURF-1 staining also was present in some of the nuclei that contained GMEB-1–GFP (b). Note, MURF-1 is also detected at the M-line region in the same myocytes (b, double arrows). Expression of GMEB-1–GFP in cardiac myocytes does not appear to affect the integrity of the COOH-terminal region of titin (d, staining with anti-titin A168–170 antibodies). Double arrows mark regular, striated titin A168–170 staining. N, nuclei. Bars, 10 μm.
and a target protein (Desterro et al., 1997; Johnson et al., 1997; Tatham et al., 2001). ISOT-3 is a member of an enzyme family responsible for cleaving isopeptide bonds. It has been proposed that Ubc9 and isopeptidases regulate the dynamics of SUMO modification (for review see Melchior, 2000; Muller et al., 2001). Further Y2H studies revealed that MURF-3 also interacts with Ubc9 (Fig. 9 A).

Interestingly, another MURF-1 binding partner that we identified through Y2H screens is GMEB-1 (Fig. 9 A). GMEB-1 is a nuclear protein that regulates transcription in response to changes in cellular glucocorticoid levels (Zeng et al., 1998, 2000b; Jimenez-Lara et al., 2000). Remarkably, GMEB-1 binding to MURF-2 or -3 was not detected in the Y2H system (Fig. 9 A), although these proteins are highly homologous to MURF-1. The interaction of MURF-1 with GMEB-1 was confirmed by glutathione-S-transferase (GST) pull-down assays under stringent conditions (Fig. 9 C). Previously, GMEB-1 mRNA was found in skeletal muscle but not in cardiac tissue by RT-PCR studies (Zeng et al., 2000a). However, it was detected in heart by Northern and Western blot analysis (Theriault et al., 1999; Jimenez-Lara et al., 2000). We performed RT-PCR analysis on heart and skeletal cDNA and detected GMEB-1 mRNA transcripts in both tissues (Fig. 9 B). (The discrepancy between our study and the Zeng et al., 2000a study might be due to differences in PCR conditions.) GMEB-1-GFP fusion proteins localized to nuclei in transfected myocytes (Fig. 9 D, c), consistent with previous studies in COS and HeLa cells (Theriault et al., 1999; Jimenez-Lara et al., 2000). In a few myocytes, GMEB-1 also was detected diffusely throughout the cytoplasm (Fig. 9 D, c), but assembly at the M-line region of the sarcomere was never observed. Cysteine pro- teins expressing GMEB-1-GFP with anti–MURF-1 antibodies demonstrated that GMEB-1-GFP and MURF-1 were both present in the same nuclei of some myocytes (Fig. 9 D, a and b), which also supports their potential interaction. Finally, cells expressing GMEB-1-GFP exhibited regular striated staining patterns for titin, thin filament, thick filament, and Z-line components, indicating that the overall sarcomeric integrity in the GMEB-1-transfected cells was not affected (Fig. 9 D, d; data shown for COOH-terminal titin staining). These data suggest that MURF-1 specifically interacts with GMEB-1. However, unlike exogenous expression of MURF-1, exogenous expression of GMEB-1 does not affect the integrity of the M-line region of cardiac sarcomeres.

Discussion

In recent years, there has been significant progress in deciphering the structure and functions of the giant sarcomeric protein titin. It appears that titin, the only known protein to span the entire half sarcomere, has multiple cellular roles. Titin contains elastic elements in its I-band region responsible for passive tension generated upon stretch; thus it functions as a molecular spring to maintain the structural integrity of contracting myofibrils. Additionally, titin may act as a "molecular blueprint" to orchestrate the assembly and organization of the thick filaments as well as other structural and regulatory components of sarcomeres. The presence of a unique Ser/Thr kinase domain at the COOH-terminal end of titin also suggests that it may participate in signal transduction pathways (for reviews with original citations see Labbé et al., 1997; Gregorio et al., 1999; Trinick and Tkshkovrova, 1999; Gregorio and Antin, 2000).

To decipher the roles of titin in sarcomeric structure, recent studies have focused on dissecting the properties of individual titin regions and their potential ligands (for reviews see McElhinny et al., 2000; Sanger and Sanger, 2001). Here, we aimed to investigate the functional significance of the interaction of titin’s COOH-terminal Ig domains A168–170 (located directly NH₂-terminal to the titin Ser/Thr kinase domain) with MURF-1, a RING finger protein. Expression of MURF-1 or titin A168–170 in primary cultures of embryonic chick cardiac myocytes severely disrupted the integrity of titin’s M-line region. The region of MURF-1 that is responsible for this phenotype was mapped to its central region, which previously has been shown to contain its titin-binding site (Centner et al., 2001). The most plausible explanation for our observations is that a dominant-negative phenotype occurred. That is, upon expression of the central region of MURF-1 or titin A168–170, the fusion proteins likely interfered with the interaction of endogenous MURF-1 with titin. Surprisingly, expression of the full-length MURF-1 molecule also resulted in this striking phenotype, suggesting that endogenous MURF-1 levels are tightly regulated. Our data suggest that MURF-1 and its interaction with titin A168–170 are critical for maintaining the stability of titin’s COOH-terminal region.

In turn, it appears that the interaction of titin with MURF-1 plays a critical role in maintaining the stability of the thick filaments. Although all of the sarcomeric components comprising the M-line region have not been elucidated, the thick filaments appear to be laterally associated with titin via their interactions with MyBP-C (along the A-band), and myomesin (at the M-line) (Houmeida et al., 1995; Obermann et al., 1997). In fact, it has been proposed that titin specifies the number and location of thick filament components (Whiting et al., 1989; Trinick, 1994; Houmeida et al., 1995). It is striking that expression of titin A168–170 in cardiac myocytes perturbed M-line titin and thick filament structure, yet expression of other titin COOH-terminal titin staining. These data suggest that MURF-1 specifically interacts with GMEB-1. However, unlike exogenous expression of MURF-1, exogenous expression of GMEB-1 does not affect the integrity of the M-line region of cardiac sarcomeres.
that certain regions of the titin filament can be “selectively” perturbed. Previous studies have shown that titin interacts with various Z-line components (α-actinin and T-cap/telethonin) and thin filament components (actin) (Funatsu et al., 1993; Jin, 1995; Granzier et al., 1997; Linke et al., 1997; Sorimachi et al., 1997; Gregorio et al., 1998; Mues et al., 1998; Young et al., 1998). From our studies, it appears that these associations along the titin molecule stabilize its NH2-terminal and I-band regions even when the structure of its COOH-terminal end is perturbed. Consistent with this idea, the NH2-terminal end of titin becomes organized during myofibril assembly before the COOH-terminal end of titin and other M-line components (Fürst et al., 1989; Schultheiss et al., 1990; Komiyama et al., 1993; van der Loop et al., 1996; Ehler et al., 1999; Rudy et al., 2001). A possible explanation for this observation is that the less organized M-line region of titin filaments may not be concentrated enough for a signal to be detected by immunofluorescence microscopy (Ehler et al., 1999). Therefore, in our study, it is likely that titin’s COOH-terminal region lost its stable interactions and was “less organized,” whereas the more NH2-terminal regions remained “bolted” to other Z- and I-band components. Interestingly, another study has demonstrated that the thick filaments remain intact when the thin filaments are perturbed upon expression of titin’s I-band, N2B region (Linke et al., 1999), in the same cell type used in our studies. Thus, titin also may function to keep the thick filaments aligned in the absence of thin filaments.

In addition to sarcomeric M-line localization, endogenous MURF-1 was detected in the nuclei of cardiac myocytes. The observation that MURF-1 assembles in only some myofibrils also indicates that its cellular levels are tightly regulated. An excellent candidate for regulating MURF-1 levels, its localization pattern, and nuclear import is SUMO-3. SUMO-3 binds to MURF-1’s RING domain (Dai and Liew, 2001), the region of MURF-1 that appears to be involved in its nuclear localization (this study). We found that two enzymes potentially involved in regulating the conjugation of SUMO with its target proteins, Ubc9 and ISOT-3, interact with MURF-1 and other MURF family members. Although the biochemical pathways and classes of enzymes involved in “SUMO modification” are parallel to those in ubiquitination, the two processes may be functionally distinct. In fact, SUMO modification has been implicated in regulating the levels and localization patterns of target proteins, including nuclear localization (for review see Melchior, 2000). All three MURF proteins exhibit multiple cellular localization patterns (Spencer et al., 2000; Centner et al., 2001; unpublished data), consistent with them being potential SUMO targets. Interestingly, SUMO-3 has recently been shown to be a key component of a new class of acute and reversible cellular stress response pathways (Saitoh and Hinchey, 2000). Further studies are needed to determine whether MURF-1 levels and/or localization patterns change in response to cellular stress, and whether titin plays a role in sensing stress response pathways in cardiac myocytes.

Strongly implicating MURF-1 with nuclear functions are the data demonstrating its specific interaction with GMEB-1 and their colocalization in the nuclei of some myocytes. GMEB-1 was first characterized as a component of a complex that binds to the GME in liver cells, thereby modulating transcription in response to glucocorticoid levels (Zeng et al., 1998; Theriault et al., 1999). Recent studies indicate that GMEB-1 is expressed in a wide range of cell types and may have roles in development and differentiation (Zeng et al., 2000a). We confirmed that GMEB-1 mRNA transcripts are present in striated muscle tissue and found that GMEB-1–GFP fusion proteins targeted to the nuclei of transfected cardiac myocytes. Given that MURF-1 is involved both in sarcomeric M-line structure (via its interaction with titin) and potentially in gene expression (via its interaction with GMEB-1), we speculate that MURF-1 acts as a link between gene expression and myofibril signaling pathways. Currently, it remains unclear what factors might modulate the MURF-1–based myofibril-to-nuclear signaling pathways. One idea is that the titin kinase domain could play a role in this process, because the MURF-1 binding site is located adjacent to this domain (Centner et al., 2001).

In conclusion, we demonstrated that the central region of MURF-1 targets to the M-line region and participates in its structural integrity in cardiac myocytes, most likely through its interaction with titin’s COOH-terminal A168–170 region. Interestingly, MURF-1 is an RBCC protein, a family whose members have been referred to as “builders of master scaffolds” because many are involved in the formation of multiprotein complexes (for review see Borden, 2000). Thus, perhaps additional factors, such as other MURF family members, are involved in regulating M-line and thick filament structure through their interactions with MURF-1. It is also striking that RING finger and RBCC proteins have been implicated in ubiquitination pathways (for review see Freemont, 2000). Consistent with this, recombinant MURF-1 protein was recently reported to have ubiquitin ligase activity (Bodine et al., 2001). Moreover, MURF-1−/− mice were resistant to skeletal muscle atrophy, suggesting that MURF-1 regulates the degradation of critical muscle proteins (Bodine et al., 2001). These results and the results from our study support our hypothesis that MURF-1 is involved in a novel pathway responsible for titin structure and/or turnover. Intriguingly, because MURF-1 also is a nuclear component that binds to the transcriptional modulator GMEB-1, MURF-1 may participate in both the regulation of myofibril assembly and structure as well as muscle gene expression. In support of our hypothesis, rat skeletal muscle MURF-1 mRNA levels increased 10-fold in response to glucocorticoid exposure (Bodine et al., 2001), which regulates GMEB-1 transcriptional activity (Theriault et al., 1999; Zeng et al., 2000b). Future studies are required to elucidate whether the dual localization of MURF-1 to myofibrils and nuclei is a result of dynamic SUMO modification.

Materials and methods

Y2H interaction studies

For a survey of potential MURF-1 interactions, a full-length MURF-1 cDNA fragment was amplified from human skeletal muscle cDNA by PCR (Sakiki et al., 1985) and inserted into pAS2-1 (Matchmaker system II; CLONTECH Laboratories, Inc.) to obtain a GAL4–BD fusion. For screening, bait constructs were transformed into Saccharomyces cerevisiae strain CG1945, P69–4A (pAS2-1 bait), or 4L0C (BTM117c bait). For some screens, the pAS2C-1 vector also was used (provided by T. Maeda, University of Tokyo). Subsequently, the cells were transformed with a human...
skeletal muscle cDNA library into the pGAD10 prey vector (HL4010A; CLONTECH Laboratories, Inc.), essentially as described by the manufacturer. Cells were plated onto SD/Leu–/Trp–/His– plates and incubated at 30°C until colonies appeared (after ~5 d). In screens where the pAS2-1 bait vector was used, the plates were supplemented with 1.5 mM 3-amino-1,2,4-triazole (Sigma-Aldrich). Transformants were picked, restreaked onto SD/Leu–/Trp–/His– plates, and prey clones were recovered by electroporation of yeast DNA in cloning buffer (2 mM Tris, 150 mM NaCl, 100 mM MgCl2) into BL21-DE3 (CLONTECH Laboratories, Inc.) cells. Whole-cell lysates in cloning buffer were prepared and subcloned into a modified pET9D vector. The constructs were transformed into Escherichia coli DH5α. His6-GST double-tagged fusion proteins were obtained by cloning into a modified pET9D vector. The constructs were transformed into BL21-DE3 (CLONTECH Laboratories, Inc.) cells. Whole-cell lysates in cloning buffer (20 mM Tris, pH 7.4, 100 mM KC1, 1% Triton X-100, plus protease inhibitors) were incubated with 20 μl of beads coated with 20 μg of bound GST fusion proteins in 300 μl binding buffer. The mixture was incubated for 1 h at 4°C, washed three times with binding buffer, and resuspended in SDS sample buffer. The protein complexes were fractionated by SDS-PAGE using 15% gels. The gels were fixed (20% methanol, 10% acetic acid), stained with Coomassie blue, treated with Amplify (Amersham Pharmacia Biotech), and photographed using a Gel Doc 2000 (Bio-Rad). The bands were quantified using image analysis software (Vinci). The ratios of each band were used to calculate the relative protein expression. The experiments were performed at least three times, and the relative protein expression was determined by normalizing the band intensity of each protein to the intensity of the 23S ribosomal RNA band. The data are presented as the mean ± standard deviation.

In vitro translation and GST pull-down experiments

In vitro translation and GST pull-down experiments were performed in the presence of [35S]methionine (Amersham Pharmacia Biotech) using a TNT T7–coupled reticulocyte lysate system according to the manufacturer’s instructions (Promega). His6-GST double-tagged fusion proteins were obtained by incubating with a modified pET9D vector. The constructs were transformed into BL21-DE3 (CLONTECH Laboratories, Inc.) cells. Whole-cell lysates in cloning buffer (20 mM Tris, pH 7.4, 100 mM KC1, 1% Triton X-100, plus protease inhibitors) were incubated with 20 μl of beads coated with 20 μg of bound GST fusion proteins in 300 μl binding buffer. The mixture was incubated for 1 h at 4°C, washed three times with binding buffer, and resuspended in SDS sample buffer. The protein complexes were fractionated by SDS-PAGE using 15% gels. The gels were fixed (20% methanol, 10% acetic acid), stained with Coomassie blue, treated with Amplify (Amersham Pharmacia Biotech), dried, and exposed using BioMax MR-1 film (Eastman Kodak Co.). The results of the Coomassie blue staining confirmed that equal amounts of each GST fusion protein were bound to the beads in the different samples (unpublished data).

Cell culture and transfection procedures

For myocyte expression studies, cDNAs containing the entire open reading frame of MURF-1 (residues 1–565) and subfragments of MURF-1, corresponding to its amino-terminal domain, were amplified from contaminating genomic DNA, RNA also was incubated with random hexamer primers and SuperScript reverse transcriptase according to the manufacturer (Strategene). To ensure that PCR products were not amplified from contaminating genomic DNA, RNA was also incubated without SuperScript reverse transcriptase. PCR amplifications were performed essentially as described by Centner et al. (2000), using MURF-1–specific primers designed to amplify 311 bp product corresponding to the MURF-1 coding region. The primers used to generate the MURF-1 fragments were: 73419S/74402R for A168–A170, 74254S/75345R for titin kinase, 74397S/75242R for mutated actin binding site, A168–A170, the COOH-terminally located kinase domain, and M-line Ig domains were amplified from total human cDNA and cloned into myosin light chain vectors. The recombinant pEGFP-C1 constructs were purified using Qiagen columns (QIAGEN) before transfection into myocytes. Plasmids were verified by sequencing. To rule out any potential artifacts resulting from the GFP tag, pCMVmyc-MURF1 constructs, as well as constructs encoding MURF-1 with the GFP tag at its COOH-terminal end, were generated (Gregorio et al., 1998) and transfected into cardiac myocytes. Identical results were obtained (unpublished data).

Cardiac myocytes were prepared from 6-d embryonic chick hearts and cultured on coverslips at 37°C. Similarly, transfected cardiac myocytes were isolated and maintained as previously described (Gustafson et al., 1987). Transfected cardiac myocytes were essentially stained as described by Gregorio et al. (1998). Cells were fixed in 2% formaldehyde–PBS for 10 min, washed in PBS, and permeabilized in 0.2% Triton X-100–PBS for 15 min. Coverslips were washed and stored in PBS at 4°C until staining. Over 200 transfected cells per construct were analyzed. Our transfection efficiencies ranged from 10–40%. All tissue culture reagents (except where noted) were purchased from Life Technologies.

Indirect immunofluorescence microscopy

Primary cultures of rat cardiac myocytes were isolated and maintained as previously described (Gustafson et al., 1987). Transfected cardiac myocytes were essentially stained as described by Gregorio et al. (1998). Cells were fixed in 2% formaldehyde–PBS for 10 min, washed in PBS, and permeabilized in 0.2% Triton X-100–PBS for 15 min. The coverslips were incubated in 2% BSA, 1% normal donkey serum–PBS for 1 h to minimize nonspecific binding of antibodies. For double-labeling experiments, cells were incubated with affinity-purified rabbit polyclonal antibodies specific to MURF-1 (5–10 μg/ml) (Centner et al., 2001), followed by Cy2-conjugated goat anti-rabbit IgG (1:600), and incubated with monoclonal sarcomeric anti-α-actinin antibodies (1:1,500) (EA-53; Sigma-Aldrich) followed by goat anti-mouse Texas red–conjugated IgG (1:600). For staining GFP-transfected cells, well-characterized antibodies were used against various sarcomeric components, including monoclonal anti-titin T1 antibodies (1:1,000; Sigma-Aldrich), monoclonal anti-myosin antibodies F59 (1:10; provided by F. Stockdale, Stanford University, Stanford, CA), rabbit anti-titin M-line–specific antibodies (1:100; Centner et al., 2001), anti-titin N2A antibodies (10 μg/ml; Centner et al., 2000), rabbit anti–MyBP-C antibodies (1:50; Linke et al., 1999), monoclonal anti-titin ABS (1:3 of cultured supernatant) provided by John Mudry, University of Leuven, Leuven, UK, and binding site antibodies 1:50 of cultured supernatant; provided by J.C. Perriard, Swiss Federal Institute of Technology, Zurich, Switzerland (Grove et al., 1984), and Texas red–conjugated phalloidin (1:200; Molecular Probes Inc.). The staining was followed by incubation with Texas red–conjugated goat anti–mouse IgG plus IgM (1:600) or Texas red–conjugated donkey anti–rabbit antibodies (1:700) for 45 min after rinse to identify nuclei. In some experiments, fixed and incubated in a DAPI stain (10 μg/ml; Sigma-Aldrich) for 5 min at room temperature before the final washing steps. Note, because of the intense diffuse- ness of soluble titin–GFP fusion proteins in transfected cells, we extracted soluble proteins with cytoskeletal (myofibril) stabilization buffer before fixation in some experiments (Gregorio and Fowler, 1995). For triple-labeling studies, a cascade blue–conjugated secondary antibody was used (1:200). All coverslips were mounted on slides using Aqua Poly/Mount (Polysciences, Inc.) and subsequently analyzed on a Zeiss Axiovert microscope using a 100× [NA 1.3] objective, and micrographs were recorded as digital images on a SenSys cooled HCCD (Photometrics). For triple-labeling studies, transfected cells were analyzed on a DeltaVision Deconvolution Model D-OL Olympus microscope with a 60× objective (1.4 NA) using a Photometrics Series 300 CCD cameras (Applied Precision). Images were processed for presentation using Adobe Photoshop 6.0. All secondary antibodies were purchased from Jackson ImmunoResearch Laboratories, except the Cy2-conjugated antibodies, which were purchased from Pierce Chemical Co.

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