Z-disc-associated, Alternatively Spliced, PDZ Motif-containing Protein (ZASP) Mutations in the Actin-binding Domain Cause Disruption of Skeletal Muscle Actin Filaments in Myofibrillar Myopathy

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Background: The binding partners of the ZASP internal region that is mutated in zaspopathy are not yet known.

Results: The internal region of ZASP binds to skeletal muscle α-actin, and zaspopathy mutations cause actin disruption.

Conclusion: ZASP mutations in the actin-binding domain are deleterious to the muscle Z-disc structure.

Significance: ZASP-actin interaction expands the role of ZASP and defines the mechanism of zaspopathy.

The core of skeletal muscle Z-discs consists of actin filaments from adjacent sarcomeres that are cross-linked by α-actinin homodimers. Z-disc-associated, alternatively spliced, PDZ motif-containing protein (ZASP)/Cypher interacts with α-actinin, myotilin, and other Z-disc proteins via the PDZ domain. However, these interactions are not sufficient to maintain the Z-disc structure. We show that ZASP directly interacts with skeletal actin filaments. The actin-binding domain is between the modular PDZ and LIM domains. This ZASP region is alternatively spliced so that each isoform has unique actin-binding domains. All ZASP isoforms contain the exon 6-encoded ZASP-like motif that is mutated in zaspopathy, a myofibrillar myopathy (MFM), whereas the exon 8–11 junction-encoded peptide is exclusive to the postnatal long ZASP isoform (ZASP-Lex10). MFM is characterized by disruption of skeletal muscle Z-discs and accumulation of myofibrillar degradation products. Wild-type and mutant ZASP interact with α-actin, α-actinin, and myotilin. Expression of mutant, but not wild-type, ZASP leads to Z-disc disruption and F-actin accumulation in mouse skeletal muscle, as in MFM. Mutations in the actin-binding domain of ZASP-LEx10, but not other isoforms, cause disruption of the actin cytoskeleton in muscle cells. These isoform-specific mutation effects highlight the essential role of the ZASP-Lex10 isoform in F-actin organization. Our results show that MFM-associated ZASP mutations in the actin-binding domain have deleterious effects on the core structure of the Z-discs in skeletal muscle.

The Z-disc of striated muscle is a highly ordered assembly of multiple proteins that defines the lateral boundary of the sarcomere, the basic contractile unit in striated muscle. The core of the Z-disc consists of an intricate lattice of actin filaments from adjoining sarcomeres that are cross-linked by α-actinin homodimers (1, 2). The Z-disc plays a critical role in the structural integrity of the sarcomere, in the transmission of force during contraction, and, likely, in mechanosensing and other cell signaling pathways (3, 4). The importance of the Z-disc in striated muscle is underscored by the fact that, in the past decade, it has become a hot spot for newly discovered genes in muscular dystrophies and cardiomyopathies, particularly the myofibrillar myopathies (MFM) (4). MFM is a descriptive term used for myopathies that are characterized by disruption of the Z-discs and accumulation of myofibrillar degradation products (5, 6). The underlying disease mechanisms for most of these myopathies have yet to be defined.

Zaspopathy is a well characterized autosomal dominant MFM (7, 8) caused by single amino acid substitutions in the Z-disc-associated, alternatively spliced, PDZ motif-containing protein (ZASP) (9). The A165V mutation is located within an evolutionarily conserved, 26-amino acid, ZASP-like motif encoded by exon 6, referred to here as striated muscle ZM or sZM. This mutation has been reported in affected members in multiple families in the United States and Europe with a consistent phenotype of MFM (9, 10). Another ZASP mutation that also is located at the sZM, A147T, causes a phenotype similar to zaspopathy and has been reported in multiple families (10).

ZASP, also known as Cypher/LDB3/Oracle (11–13), is a member of the PDZ-LIM domain family and plays an important role in maintaining the structural integrity of the striated muscle Z-discs in multiple species, including mouse and Drosophila (14, 15). ZASP has six alternatively spliced isoforms that are cardiac- or skeletal muscle-specific in mouse and human (16, 17). The Z-disc associated, alternatively spliced, PDZ motif-containing protein (ZASP) interacts with homodimers. Z-disc-associated, alternatively spliced, PDZ in skeletal muscle.
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17). In human skeletal muscle, alternative splicing of exons 9 and 10 generates three isoforms (Fig. 1, inset) (10, 16). The long isoforms, either containing exon 10 (ZASP-L) or lacking exon 10 (ZASP-LΔex10), include an N-terminal PDZ domain, an internal sZM, and three C-terminal LIM domains. ZASP-L is replaced by ZASP-LΔex10 in postnatal skeletal muscle (17). The short isoform (ZASP-S) has a stop codon in exon 9 and lacks the LIM domains. The PDZ domain encoded by exons 1–3 interacts with the C terminus of α-actinin-2, a structural component of the Z-disc, and other Z-disc-associated proteins, such as myotilin and calsarcin 1 (11, 12, 14, 18, 19). The LIM domains encoded by exons 12–16 bind to protein kinase C and, potentially, have a role in protein kinase C-mediated signaling pathways (11). However, proteins that interact with the internal region of ZASP have not yet been identified in skeletal muscle.

EXPERIMENTAL PROCEDURES

DNA Constructs—The full-length coding sequence of ZASP-S (GenBank accession no. BC010929) and its fragment encoding amino acids 108–239 (sZM-132 aa) were amplified by PCR with human LDL3 cDNA as a template (IMAGE4291498, Open Biosystems). The full-length coding sequence of ZASP-L was obtained in two steps. A fragment encoding exons 1–7 was amplified by PCR with ZASP-S as a template and cloned into vector pcDNA3. Subsequently, a fragment encoding exons 7–16 (without exon 9) was amplified with human LDL3 cDNA as a template (IMAGE40080656, K. K. Dnaflow, Yokohama City, Japan) and added to the exon 1–7 LDL3 clone with a unique EcoRI restriction site in exon 7 to obtain a full-length ZASP-L construct. Fragments encoding human LDL3 exon 6 and exons 8–10–11 were amplified by PCR with ZASP-L as a template. A fragment of ZASP cDNA with deletion of either sZM or exon 10 was generated by Gene Synthesis (GenScript) and incorporated into ZASP constructs by a fragment swap using unique BstEII, EcoRI, and Bsu36I restriction sites within the template. A fragment of ZASP cDNA with deletion of either sZM or exon 10 was amplified by PCR with ZASP-S as a template and cloned into vector pcDNA3. Subsequently, a fragment encoding exons 7–16 (without exon 9) was amplified with human LDL3 cDNA as a template (IMAGE40080656, K. K. Dnaflow, Yokohama City, Japan) and added to the exon 1–7 LDL3 clone with a unique EcoRI restriction site in exon 7 to obtain a full-length ZASP-L construct. Fragments encoding human LDL3 exon 6 and exons 8–10–11 were amplified by PCR with ZASP-L as a template. A fragment of ZASP cDNA with deletion of either sZM or exon 10 was generated by Gene Synthesis (GenScript) and incorporated into ZASP constructs by a fragment swap using unique BstEII, EcoRI, and Bsu36I restriction sites within the cDNA. The A165V and the A147T mutations were introduced by site-directed mutagenesis. These ZASP cDNA fragments and full-length constructs were cloned into the Y2Hbait vector as a template (IMAGE4291498, Open Biosystems). The short isoform (ZASP-S) has a stop codon in exon 9 and lacks the LIM domains. The PDZ domain encoded by exons 1–3 interacts with the C terminus of α-actinin-2, a structural component of the Z-disc, and other Z-disc-associated proteins, such as myotilin and calsarcin 1 (11, 12, 14, 18, 19). The LIM domains encoded by exons 12–16 bind to protein kinase C and, potentially, have a role in protein kinase C-mediated signaling pathways (11). However, proteins that interact with the internal region of ZASP have not yet been identified in skeletal muscle.

Antibodies—The following primary antibodies were used: mouse anti-ZASP (catalog no. H00011155-M06, Abnova), rabbit anti-α-actinin-2 (catalog no. 2310-1, Epitomics), rabbit anti-myotilin (catalog no. ab68915, Abcam), mouse anti-α-tubulin (catalog no. T6199, Sigma), rabbit anti-HA tag (catalog no. ab9110, Abcam), mouse anti-FLAG tag (catalog no. F1804, Sigma), mouse anti-skeletal muscle α-actin (catalog no. 5C5, Sigma), mouse anti-GST (catalog no. G1160, Sigma), and goat anti-GST (catalog no. 27-4577-01, GE Life Sciences).

Yeast Two-hybrid Screening—A yeast two-hybrid screen was performed using the Matchmaker Gold system (Clontech). Briefly, Y2HGold yeast cells were transformed with the plasmid pGBK7 encoding the GAL4 DNA binding domain fused in frame to sZM-132aa WT or A165V. Transformants were mated with Y187 yeast containing the plasmid pGADT7 encoding cDNAs from a human skeletal muscle cDNA library fused to the GAL4 transcription activation domain. Diploids were plated onto quadruple drop-out nutrition selection plates (-Leu/Trp/His/Ade) containing Aureobasidin A and X-α-Gal. The colonies were put through two rounds of isolation. Prey-containing pGADT7 plasmids were amplified in Escherichia coli, and the inserts were sequenced.

Yeast Two-hybrid Pairwise Assay—Yeast two-hybrid pairwise experiments were performed using the Matchmaker Gold system (Clontech). The experiments were carried out with the mating strategy described in the Clontech Yeast Protocols Handbook, with the bait and prey constructs in the Y2HGold strain. The activity of the nutritional reporter genes was assayed by culturing on different selection plates (S.D.-Leu/-Trp, S.D.-Leu/-Trp/-His, and S.D.-Leu/-Trp/-His/Ado). The pair pGBK7–53/pGADT7-T served as a positive control in all experiments. As negative controls, appropriate empty vectors tested against the different bait and prey constructs.

Immunoprecipitation—Mouse vastus muscles were homogenized on ice in 20 mM Tris (pH 7.2), 100 mM KCl, 5 mM EGTA, 1% Triton X-100, and complete protease inhibitor mix (Roche Applied Science), incubated for 1 h at 4 °C with gentle agitation, and then centrifuged at 3000 × g for 30 min at 4 °C. The supernatants were precleared with protein A Dynabeads (Invitrogen) for 1 h at 4 °C. The cleared muscle lysates were incubated with either the mouse anti-ZASP monoclonal antibody or normal mouse IgG (Santa Cruz Biotechnology) for 3 h and subsequently combined with protein A Dynabeads (Invitrogen) for an overnight rotation at 4 °C. Transfected COS7 or HEK293 cells were sonicated in ice-cold lysis buffer (20 mM Tris-HCl (pH 7.2), 5 mM EGTA, 100 mM KCl, 1% Triton X-100, 1 mM DTT, and complete protease inhibitor mix). Protein G Dynabeads (Invitrogen) were incubated with either mouse anti-FLAG tag antibody or normal mouse IgG, combined to cleared cell lysates, and rotated for 3 h at 4 °C. The beads retaining the immune complexes were washed three to five times either with PBS/0.02% Tween 20 or with a buffer containing 50 mM Tris-HCl (pH 8.0), 500 mM NaCl, 0.1% SDS, 1% Triton X-100, 5 mM EDTA, 5 mM EGTA, and complete protease inhibitor mix and
then heated in SDS sample loading buffer for 5 min. The eluates and total lysates were analyzed by immunoblotting.

Protein Electrophoresis and Immunoblotting—Protein samples were resolved on 10% Novex Tris-glycine gels (Invitrogen) and then detected either by staining with colloidal blue (Invitrogen) or immunoblotting. For immunoblots, the proteins were transferred to a nitrocellulose membrane (Invitrogen). After blocking in 5% milk in PBS/0.05% Tween 20 or PBS/1% BSA for 1 h at room temperature, the membranes were incubated with primary antibody either overnight at 4 °C or for 1 h at room temperature and then with secondary antibodies for 1 h at room temperature. Secondary antibodies were conjugated either with HRP (Jackson ImmunoResearch Laboratories) or IRDye (Li-Cor Biosciences). Either the Chemidoc XRS (Bio-Rad) or Odyssey (Li-Cor) system was used for detection.

Purification of Wild-type and Mutant ZASP from E. coli—Cells expressing GST-ZASP-LΔex10 or His6-ZASP-S proteins were disrupted with a French press in 25 mM Tris-HCl (pH 8.0) plus 500 mM NaCl (GST-ZASP-LΔex10) or PBS buffer plus 0.2% Tween 20 (His6-ZASP-S). For GST-tagged ZASP-LΔex10 proteins, the clarified cell extract after centrifugation was filtered with a 0.45-μm filter and combined into glutathione-Sepharose 4 Fast Flow Beads resins (GE Life Sciences). The proteins were eluted using the protocol of the manufacturer. Eluted protein samples were run through a Superdex 200 10/300 gel filtration column (GE Life Sciences) connected to an AKTA Explorer FPLC system (Amersham Biosciences/GE Life Sciences). The sample volume injected for each experiment was 0.5 ml. The elution rate was 0.3 ml/min, and the volume per fraction was 0.5 ml. The buffer composition was 10 mM Tris-HCl (pH 8.0), 50 mM NaCl, and 10% glycerol (storage buffer). Each protein was eluted in an 11-ml final volume. The purified GST-ZASP proteins were further concentrated in the storage buffer using Amicon Ultra-4 centrifugal filter units (Millipore). The working aliquots of proteins were snap-frozen in liquid nitrogen and stored at −70 °C. For His6-tagged ZASP-S proteins, expression resulted in accumulation of mostly insoluble protein, which was extracted with 8 mM guanidine-HCl and then diluted with an equal volume of 4 mM guanidine-HCl. The extract was clarified by centrifugation. The solubilized and unfolded proteins were purified by nickel affinity using Streamline chelating resin (GE Healthcare), followed by gel filtration using Superdex S75. All buffers included 4 mM guanidine-HCl. Proteins were then folded (1 mg/ml) by sequential dialysis against 50 mM sodium acetate (pH 5.0) containing 3–0 M urea, followed by buffer exchange to 25 mM sodium phosphate (pH 6.1). Any aggregated protein was removed by centrifugation at 150,000 g for 1 h, followed by filtration using 0.22-μm Millex-GV (Millipore). The purified His6-ZASP-S proteins were concentrated using Amicon Ultra-4 centrifugal filter units (Millipore) and stored in 25 mM sodium phosphate buffer (pH 6.1) at 4 °C.

GST Pulldown Assay—Mice vastus muscles were homogenized on ice in 50 mM Tris-HCl (pH 7.5), 150 mM NaCl, 1% Nonidet P-40, 2 mM EGTA, 2 mM EDTA, 5% glycerol, 1 mM DTT, and EDTA-free complete protease inhibitor mix, incubated for 1 h at 4 °C with gentle agitation, and then centrifuged at 3000 × g for 30 min at 4 °C. The supernatants were precleared with glutathione magnetic beads (Pierce) for 10 min at 4 °C. The cleared muscle lysates were incubated with GST-ZASP-LΔex10 (wild-type, A165V, and A147T) and GST-bound to glutathione magnetic beads for 3 h at 4 °C. The beads retaining the protein complexes were washed five times with 50 mM Tris-HCl (pH 8), 500 mM NaCl, 0.1% SDS, 1% Triton X-100, 5 mM EGTA, 5 mM EDTA, and complete protease inhibitor mix and heated in SDS sample loading buffer for 5 min. The eluates and muscle lysates were analyzed by immunoblotting. Multiple independent experiments were performed (n = 4) using vastus muscle lysates from three wild-type mice.

Slot Blot Overlay Assay—Equal amounts of GST and GST-ZASP-LΔex10 proteins (120 pmol) were diluted in buffer (5 mM Tris-HCl (pH 8.0) and 0.2 mM CaCl2) and loaded on a nitrocellulose membrane using a Bio-Dot SF microfiltration apparatus (Bio-Rad). After blocking in 5 mM Tris-HCl (pH 8.0), 0.2 mM CaCl2, 1 mM DTT, and 2% BSA for 1 h, the membrane was incubated with biotinylated G-actin (46.5 nM, Cytoskeleton, Inc.) in blocking buffer containing 0.2 mM ATP overnight at 4 °C and then washed with 50 mM Tris-HCl (pH 8.0), 150 mM NaCl, 1 mM DTT, and 0.1% Tween 20 for 30 min at room temperature. The membrane was incubated with HRP-conjugated streptavidin (N100, Thermo Scientific), and G-actin binding was detected using chemiluminescence. GST and GST-ZASP-LΔex10 proteins were detected by immunoblotting. The Arp2/3 protein complex and BSA were used as controls in this assay. Multiple independent experiments were performed (n = 3).

Actin Filaments and ZASP High-speed Cosedimentation Assay—The binding of ZASP to skeletal muscle α-actin filaments was examined with the actin binding protein spin-down assay biochem kit (Cytoskeleton, Inc.). Briefly, proteins were clarified by centrifugation at 150,000 g for 1 h at 4 °C, and the supernatants were collected. Rabbit skeletal muscle G-actin (0–16 μM, Cytoskeleton Inc.) was polymerized in 5 mM Tris-HCl (pH 7.5), 0.2 mM CaCl2, 40 mM KCl, 2 mM MgCl2, and 1 mM ATP for 1 h at room temperature in the presence of clarified GST-ZASP-LΔex10 (0.2 μM) or His6-ZASP-S (2 μM) proteins. Subsequently, samples were centrifuged at 150,000 g for 1.5 h at 24 °C to pellet F-actin and F-actin binding proteins. Comparable amounts of supernatant and pelleted fractions were analyzed by densitometric scanning of colloidal blue-stained (Invitrogen) SDS/polyacrylamide gels. The fraction of protein pelleted in the absence of F-actin was subtracted from the values obtained in the presence of F-actin. Binding affinities were determined by nonlinear regression analysis with GraphPad Prism software. Average trends from triplicate assays are shown.

Intramuscular Injections of ZASP-GFP cDNA—Wild-type mice (C57BL/6) 3 months of age were anesthetized by isoflurane inhalation. Tibialis anterior (TA) muscle was pretreated by intramuscular injection of bovine hyaluronidase (15 μl, 0.4 units/μl, Sigma-Aldrich). Two hours later, EGFP-tagged ZASP-LΔex10-WT or ZASP-A165V cDNA (75 μg) in a total volume of 25 μl of PBS was injected into the muscles of opposite limbs with a 30-gauge needle. The muscle was then electroporated using TaiChi Power 80 acupuncture needle electrodes (LhasaOMS) connected to an Electro Square Porator (ECM 830, BTX A Division of Genetronics, Inc.). Electroporation parameters were 175 V/cm, eight pulses at 1 Hz, and 20-ms duration per pulse. The mice were sacrificed, and the TA muscles were harvested for immuno-
staining or electron microscopy 1 week or 4 weeks after electroporation. The intramuscular injections and all analyses were performed by blinded investigators.

**Immunofluorescence**—C2C12 cells and frozen sections (6- to 9-μm thickness) of human vastus lateralis muscle or mouse TA muscle were fixed in 2–3% paraformaldehyde for 15 min, permeabilized with 0.2% Triton X-100 in 2.5% normal goat serum/PBS for 20 min at room temperature, and then incubated with primary antibody overnight at 4 °C followed by Dylight 488-, 594-, or 633-conjugated goat anti-mouse or anti-rabbit IgG (Jackson ImmunoResearch Laboratories) for 1 h at room temperature. Actin filaments were stained with either rhodamine- or Alexa Fluor 594-phalloidin (Invitrogen) for 1 h at room temperature. Images were acquired on a confocal microscope (TCS SP5 II (Leica) or LSM710 (Carl Zeiss)) using a ×63/numerical aperture 1.4 oil immersion objective at room temperature and processed with Adobe Photoshop CS5. Raw images were exported as TIFF files, and identical settings were used on all images of a given experiment for any adjustments in image contrast and brightness. Image analysis was performed with LAS AF (Leica) and ImageJ (National Institutes of Health) software.

**Electron Microscopy**—Mouse muscle tissue was selected from areas of TA muscle showing uniform GFP fluorescence of transfected ZASP-GFP proteins in muscle fibers. The muscle tissue blocks were fixed with 2% glutaraldehyde and 2% paraformaldehyde in 0.1 N cacodylate buffer. The embedding and staining were carried out in the NINDS, National Institutes of Health electron microscopy facility using a standard approach. Images were acquired on an electron microscope (1200EX II, Jeol, Inc.) and processed with Adobe Photoshop CS5.

**RESULTS**

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**FIGURE 1. Characterization of ZASP-skeletal muscle α-actin 1 (ACTA1) interaction in yeast cells.** Representative images of Y2H pairwise assays demonstrating the interaction of different regions of ZASP and ACTA1 are shown. Positive interactions were determined by yeast growth on the media deficient in HIS3 and ADE2. The transformation efficiency was uniform for all constructs (right panels). Sequential 10-fold yeast dilutions are shown. Inset, the human LDB3/ZASP genomic structure and the three skeletal muscle-specific ZASP splice isoforms: the pre-natal long isoform (L), the post-natal long isoform (LΔex10), and the short isoform (S). The known domains of ZASP are highlighted. The location of ZASP A147T and A165V mutations in the sZM domain is indicated. The peptide sZM-132aa, wild-type or A165V, was used as bait in a Y2H screen. A, Y2H pairwise assays show interactions between the indicated ZASP regions and the peptide encoding amino acids 247–377 in the C terminus of ACTA1. WT and mutant (A165V and A147T) sZM-132aa interact with the ACTA1 peptide. ZASP exon 6-encoded peptide, which contains the sZM domain, interacts with the C terminus of ACTA1. Yeast cells cotransformed with empty bait vector and the ACTA1 peptide prey show no growth on the interaction media (−). ZASP-S interacts with the ACTA1 peptide, and deletion of the sZM domain (ΔsZM) abolishes the interaction. The ZASP peptide encoded by exons 8–11Δex10, corresponding to ZASP-LΔex10, interacts with the ACTA1 peptide, and inclusion of exon 10, corresponding to ZASP-L, abolishes the interaction of this ZASP region with the C terminus of ACTA1. B and C, Y2H pairwise assays show the interaction between the indicated ACTA1 regions and ZASP sZM-132aa (B) and exon 8–11Δex10 (C) baits. The ZASP baits interact with amino acids 247–377 of ACTA1, corresponding to the clones identified in our Y2H screen. ACTA1 peptides encoding overlapping amino acid residues within the C terminus were tested against the ZASP baits. Deletion of ACTA1 amino acids 287–303 and 304–325 abolished the interaction with ZASP peptides encoded by exons 8–11Δex10 and the sZM-132aa (wild-type and A165V and A147T mutants), respectively.
Characterization of ZASP-skeletal muscle α-actin 1 (ACTA1) interaction in mammalian cells and in vitro with purified proteins. A, co-IP (IP) assay showing the interaction between the ZASP isoforms and ACTA1 in wild-type mouse vastus muscle lysates. B and C, co-IP assays showing the interaction between the indicated ZASP isoforms and ACTA1 in non-muscle (COS7/HEK293) cells. The proteins were detected with anti-FLAG and anti-HA antibodies. D, purified GST-tagged ZASP-LΔex10 (WT, A165V, and A147T), but not GST alone, binds to biotinylated G-actin (G-actin) in a slot blot overlay assay. Biotin-G-actin was detected with HRP-streptavidin. GST and GST-ZASP were detected by immunoblotting with GST antibody (anti-GST). Arp2/3 and BSA served as positive and negative controls for G-actin binding. E, high-speed cosedimentation assay of purified GST or the indicated GST-ZASP-L or exons 8–11 corresponding to ZASP-L and ACTA1 (247–377), and this interaction was abolished by deletion of the thin filaments in adult skeletal muscle. It is a single chain metallothionein 2A (MT2A). The C terminus of PGM1 has been identified previously as an interactor of ZASP in a Y2H screen of a human heart cDNA library (20). We validated ZASP interaction with ACTA1 as discussed below. MYBPC1 is currently under investigation. The other four proteins were regarded as probable false positives.

For ACTA1, all prey clones (n = 8 for each bait) encoded the 131 C-terminal amino acids 247–377. The interaction between sZM-132aa-WT and sZM-A165V and ACTA1 (247–377), corresponding to the original clones in the Y2H screen, was validated by a Y2H pairwise assay (Fig. 1A). The sZM-132aa-A147T bait also interacted with ACTA1 (247–377) in yeast (Fig. 1A). The ZASP peptide encoded by exon 6, which contains the sZM domain, alone was sufficient for interaction with the C terminus of ACTA1 (Fig. 1A), demonstrating that the ZASP regions encoded by exons 5 and 7 are not required for ZASP-actin interaction in yeast cells. ZASP-S interacted with ACTA1 (247–377), and this interaction was abolished by deletion of the 26 amino acids corresponding to the sZM (Fig. 1A). We tested the peptides encoded by exons 8–10–11 corresponding to ZASP-L or exons 8–11Δex10 corresponding to ZASP-LΔex10 against ACTA1 (247–377) in yeast cells. Although the peptide encoded by exons 8–11Δex10 interacted with the C terminus of skeletal α-actin, this interaction was abolished by inclusion of exon 10 (Fig. 1A). Thus, the sZM domain that is mutated in zaspopathy is important for the interaction between ZASP and α-actin in skeletal muscle. ZASP-LΔex10 contains an additional actin-binding region encoded by the exon 8–11 junction that is not present in the other ZASP isoforms.

Skeletal muscle α-actin is a highly conserved component of the thin filaments in adult skeletal muscle. It is a single chain peptide with 377 amino acids. To further define the region of α-actin that binds to ZASP, different prey clones encoding shorter fragments of ACTA1 spanning the entire coding sequence were tested against either sZM-132aa (WT and mutant) or the exon 8–11Δex10-encoded peptide of ZASP in Y2H pairwise assays. As anticipated from the Y2H screen, the ZASP peptides selectively interacted with amino acids 247–377 of skeletal α-actin (Fig. 1, B and C). Serial deletion analysis of ACTA1 (247–377) showed that the amino acids 287–325 of skeletal muscle α-actin are required for binding to ZASP in yeast cells (Fig. 1, B and C).

The binding of ZASP to α-actin in skeletal muscle was validated by coimmunoprecipitation (co-IP) of these proteins from muscle lysates of adult wild-type mice. The ZASP antibody
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FIGURE 3. Characterization of the interaction between ZASP and α-actinin-2 (ACTN2) in yeast cells. A, diagram of the ACTN2 domain structure. ACTN2 has two N-terminal actin-binding domains (ABD), an internal spectrin rod domain, and the C-terminal EF-hands. ACTN2 amino acids encoding these domains are numbered as in the human sequence and shown in parentheses below each domain. B, Y2H pairwise assays show that the internal ZM region of ALP (107–273) interacts with the spectrin rod domain of ACTN2. In the same assay, WT and mutant (A165V and A147T) sZM-132aa do not interact with the spectrin rod domain of ACTN2. C, Y2H pairwise assays show that ZASP-S (WT and A165V) interacts with the C terminus of ACTN2. This interaction persists after deletion of the sZM domain (∆sZM). Neither the wild-type nor mutant ZASP-sZM-132aa baits interact with the C terminus of ACTN2. Transformation efficiency was uniform for all constructs (right panel). Sequential 10-fold yeast dilutions are shown.

ZASP, Skeletal Muscle α-Actin, and the α-Actinin-2 Protein Complex at the Z-discs of Skeletal Muscle—Skeletal actin filaments are cross-linked by adjacent sarcomeres cross-linked by antiparallel homodimers of α-actinin-2. The predicted position of ZASP-LΔex10 between α-actinin-2 and skeletal α-actin filaments is shown. The domain structures of the proteins are as shown in A.

The actin-binding domain of ZASP containing the sZM-132aa (WT) (red) interacts with the C terminus of α-actinin-2 (green) and exon 8–11Δex10-encoded peptide (black) binds to the C terminus of α-actin, and amino acids 287–325 of α-actin (dark orange) are required for this interaction in yeast cells (this study). The ZASP A165V and A147T mutations do not disrupt these interactions. B, schematic of the skeletal muscle Z-disc core structure showing skeletal actin filaments (F-actin) from adjacent sarcomeres cross-linked by antiparallel homodimers of α-actinin-2. The predicted position of ZASP-LΔex10 between α-actinin-2 and skeletal α-actin filaments is shown. The domain structures of the proteins are as shown in A.

The actin-related protein 2/3 (Arp2/3) complex and BSA were used as positive and negative controls, respectively, in the G-actin binding assay. ZASP binding to skeletal actin filaments (F-actin) was examined by a high-speed cosedimentation assay. Wild-type and mutant versions of GST-ZASP-LΔex10 proteins bound to F-actin with micromolar affinity ($K_d$ mean ± S.E.; WT, 0.7 ± 0.25 μM; A165V, 0.9 ± 0.27 μM; and A147T, 0.8 ± 0.13 μM), whereas GST alone did not sediment with F-actin (Fig. 2E). To examine the effect of zaspopathy mutations on the exon 6-binding site, we used His$_6$-ZASP-S proteins in F-actin cosedimentation assays. We observed a similar F-actin binding affinity of the sZM domain between WT and mutant proteins ($K_d$ mean ± S.E.; WT, 0.9 ± 0.36 μM; A165V, 1.1 ± 0.31 μM; and A147T, 1.5 ± 0.39 μM).

ZASP, Actinin-associated LIM protein (ALP), and spectrin-actin-binding protein (ZASP) form a complex at the Z-discs of skeletal muscle, and this complex is disrupted in myofibrillar myopathy.
ALP also belongs to the Enigma family and localizes to skeletal muscle Z-disc by binding to α-actinin (22, 23). ALP has a 26-residue conserved sequence between the PDZ and LIM domains, which is named ZASP-like motif (ZM). The ZM-containing internal region of ALP (aa 107–273) interacts with the spectrin rod domain of α-actinin (24). Similarly, it has been proposed that ZASP-sZM-132aa interacts with the spectrin rod domain of α-actinin-2 (25). However, no experimental evidence exists for direct binding between ZASP-sZM-132aa and the actinin rod. Indeed, localization of GFP-sZM-132aa to α-actinin was dependent on the integrity of actin filaments in C2C12 cells (25). We tested the interaction of sZM-132aa with the α-actinin-2 spectrin rod domain and compared it to the binding ability of ALP (107–273) to the actinin rod domain of α-actinin-2 (24), and we could not detect interaction between ZASP-sZM-132aa and the actinin rod. Indeed, localization of GFP-sZM-132aa to α-actinin was dependent on the integrity of actin filaments in C2C12 cells (25). We tested the interaction of sZM-132aa with the α-actinin-2 spectrin rod domain, as reported previously (24), and we could not detect interaction between ZASP-sZM-132aa (wild-type or mutant) and the α-actinin-2 spectrin rod domain (Fig. 3B). The PDZ domain of ZASP interacts with the C terminus of α-actinin-2 (11, 12, 14, 18). ZASP-PDZ domain interaction with the C terminus of α-actinin-2 was not affected by the A165V mutation located in the sZM domain or by deletion of the sZM domain (Fig. 3C). ZASP-sZM-132aa did not interact with the C terminus of α-actinin-2 in yeast cells (Fig. 3C). These observations indicate that ZASP forms a protein complex with skeletal muscle α-actin and α-actinin-2 at the Z-discs of skeletal muscle via distinct binding domains, as summarized in Fig. 4.

Isoform-specific Effects of the ZASP A165V and A147T Mutations on Actin Stress Fibers in Mouse Muscle (C2C12) Cells—Actin stress fibers of cultured cells are composed of F-actin bundles that are held together by α-actinin (26). The amino acid sequence of various actins in muscle and non-muscle tissues is highly conserved (∼90%), with a majority of amino acid changes found in the N terminus region (27). We examined the effects of the ZASP A165V and A147T mutations in all three skeletal muscle-specific ZASP isoforms on actin stress fibers in muscle cells. The ZASP-GFP fusions were transfected in C2C12 myoblasts. After 48 h, the morphology of the actin cytoskeleton was examined in transfected cells that had a similar range of GFP fluorescence intensity values (range, 20–86 arbitrary units; mean ± S.E., WT 44 ± 2.4 arbitrary units, A165V 49 ± 2.4 arbitrary units, and A147T 51 ± 1.9 arbitrary units; n > 39 cells for each isoform). The GFP-tagged ZASP-S, ZASP-L, and ZASP-LΔex10 isoforms localized to actin stress fibers (Figs. 5 and 6). We observed that the expression of the ZASP-LΔex10-GFP mutants (A165V and A147T), but not the wild-type pro-
tein, consistently caused disruption of actin stress fibers in transfected muscle cells (Fig. 5). The organization of microtubules was preserved in cells with a disrupted actin cytoskeleton, demonstrating selective vulnerability of actin filament bundles to the toxic effects of the mutant ZASP-L\(^\text{H9004}\text{ex10}\) proteins. Organization of the actin cytoskeleton appeared normal in muscle cells overexpressing wild-type and mutant ZASP-S-GFP and ZASP-L-GFP proteins (Fig. 6). Similar observations were made in mouse embryonic fibroblast (NIH3T3) cells transfected with the N-terminal FLAG- or HA-tagged ZASP fusion proteins (data not shown). Thus, the disruptive effects of ZASP mutants (A165V and A147T) on actin filaments were only evident when expressing ZASP-L\(^\text{H9004}\text{ex10}\), but not other skeletal muscle-specific isoforms, in cultured cells.

Mutant ZASP-L\(^\text{H9004}\text{ex10}\) Causes Disruption of Z-discs and Sarcoplasmic Accumulation of Actin Filaments in Adult Mouse Skeletal Muscle—To examine the effects of the ZASP-L\(^\text{H9004}\text{ex10}\)-GFP mutant on F-actin in skeletal muscle, WT and A165V versions were expressed by electroporation in TA muscles of opposite limbs in 3-month-old C57BL/6J wild-type mice (\(n = 7\)). We compared the distribution of F-actin and \(\alpha\)-actinin-2, the anchor of F-actin at Z-discs, in the transfected muscle fibers expressing ZASP-L\(^\text{H9004}\text{ex10}\)-A165V-GFP (referred to here as ZASP-A165V-GFP) protein to those expressing ZASP-L\(^\text{H9004}\text{ex10}\)-WT-GFP (referred to as ZASP-WT-GFP) protein in the same animal. Longitudinal sections of the TA muscles were immunostained for \(\alpha\)-actinin-2 or incubated with phalloidin to stain F-actin 1 week and 4 weeks after electroporation. Wild-type and mutant ZASP-GFP colocalized with \(\alpha\)-actinin-2 at Z-discs in the muscle fibers (Fig. 7, A and B). The architecture of Z-discs and F-actin was preserved in the muscle fibers expressing ZASP-WT-GFP 1 week and 4 weeks after electroporation (Figs. 7–9). In contrast, we observed a loss of \(\alpha\)-actinin-2 from Z-discs in the muscle fibers expressing ZASP-A165V-GFP as early as 1 week after electroporation (Fig. 7A). By 4 weeks, there was a generalized loss of \(\alpha\)-actinin-2 from Z-discs of the transfected muscle fibers, whereas ZASP-A165V-GFP was localized to the Z-discs (Fig. 7B). The GFP fluorescence of ZASP-A165V-GFP was also seen in the sarcoplasm between the Z-discs in some areas (Fig. 7B).

We saw accumulations of phalloidin-stained F-actin that colocalized with ZASP-A165V-GFP in the sarcoplasm of many fluorescent muscle fibers (Fig. 7C). Such F-actin accumulations were never observed in the muscle fibers overexpressing ZASP-WT-GFP. Sarcoplasmic accumulations of ZASP, which colocalized with phalloidin-stained F-actin, were also detected in

![Image of ZASP-Actin Interaction in Myofibrillar Myopathy](image-url)
Previous studies reported prominent accumulations of myotilin in skeletal muscle fibers of patients with zaspopathy (9, 10). Myotilin interacts with ZASP and α-actinin and controls sarcomere assembly by cross-linking actin filaments (19, 28, 29) (Fig. 8A). Wild-type and mutant GST-tagged ZASP pulled down myotilin from mouse muscle lysates (Fig. 8B). Immunostaining of the mouse TA muscle fibers showed a loss of α-actinin-2 from the Z-discs 1 week after electroporation. 8, at 4 weeks, ZASP-A165V localizes to the Z-discs, whereas there is a generalized loss of α-actinin-2 from the Z-discs in the muscle fibers. We also observed the GFP fluorescence of ZASP-A165V in the sarcoplasm between the Z-discs. Relative fluorescence intensity distributions of ZASP-WT and ZASP-A165V and α-actinin-2 across the white lines (40 μm) in the merged images are shown in the right panels. C, phalloidin staining of mouse TA muscle fibers showing that ZASP-A165V (green) and F-actin (red) colocalize in sarcoplasmic accumulations 4 weeks after electroporation. D, immunostaining with ZASP antibody and phalloidin shows colocalization of ZASP (green) and F-actin (red) accumulations in the sarcoplasm of the muscle fibers of a patient. Scale bar = 10 μm (A and B) and 25 μm (C and D).

DISCUSSION

ZASP is required for maintaining the structure of Z-discs in skeletal muscle and heart (14, 30). ZASP interaction with α-actinin-2 and other Z-disc proteins is necessary, but not sufficient, to maintain the structural integrity of striated muscle Z-discs (31). In this study, we show that ZASP directly binds to skeletal muscle α-actin, which is fundamentally important to the Z-disc architecture in postnatal skeletal muscle (32, 33).
The actin-binding domain is between the modular protein-interacting PDZ and LIM domains. It is noteworthy that mutations associated with cardiac and skeletal myopathies are located in this region of ZASP (9, 10, 16, 25). The actin-binding sites in ZASP are also expressed in heart muscle, but the composition of cardiac transcripts that contain the binding sites is not yet known (16, 17). Up to 20% of cardiac muscle actin consists of the skeletal muscle α-actin isoform (34). The cardiac and skeletal muscle α-actin isoforms are 99% identical, with only four amino acid differences (27), and they have similar functions in postnatal skeletal muscle (35). The multiple ZASP isoforms may play unique roles in maintaining the Z-disc structure by interacting with α-actin in heart as well as skeletal muscle.

Our Y2H and co-IP studies identified two actin-binding sites that are differentially expressed in multiple ZASP isoforms. The exon 6-encoded sZM domain that is mutated in zaspopathy is common to all ZASP isoforms in skeletal muscle. In contrast, the exon 8–11 junction-encoded peptide is present exclusively in the postnatal long ZASP isoform (ZASP-LΔex10) in skeletal muscle (17). Zaspopathy mutations in ZASP-LΔex10, but not the ZASP-S and ZASP-L isoforms, cause disruption of the actin cytoskeleton. This finding highlights the essential role of the ZASP long isoform in the Z-disc structure in skeletal muscle and corroborates isoform-specific deletion studies in mice (31). Because the PDZ, sZM, and LIM domains are common to both long isoforms, the actin-binding region of ZASP-LΔex10,

FIGURE 8. Effects of ZASP-LΔex10-A165V on myotilin in mouse skeletal muscle. A, diagram of skeletal muscle Z-disc showing interaction of myotilin with ZASP, skeletal muscle α-actin, and α-actinin-2. The domain structures of the proteins are same as those shown in Fig. 4. B, GST pulldown assay showing the interaction between wild-type and mutant (A165V and A147T) GST-ZASP-LΔex10 proteins with myotilin in wild-type mouse vastus muscle lysates. C, longitudinal sections of mouse tibialis anterior muscles electroporated with WT and mutant (A165V) versions of ZASP-LΔex10-GFP. Immunostaining with myotilin antibody shows localization of myotilin to the Z-discs in muscle fibers. ZASP-WT colocalizes with myotilin at the Z-discs of transfected muscle fibers. In contrast, there is a loss of myotilin from the Z-discs of the muscle fibers expressing ZASP-A165V 4 weeks after electroporation. Focal accumulation of myotilin and ZASP-A165V are seen in the sarcoplasm of the muscle fiber with evidence for colocalization of the two proteins in some areas. Scale bars = 10 μm.
including the exon 8–11 junction-encoded peptide, likely plays an important role in mediating the abnormal actin phenotype of the A165V and A147T mutants.

Striated muscle has precise and regular arrangements of actin filaments that are organized in sarcomeres. Such a precise assembly of actin filaments by cross-linking proteins is indispensable to the structure and function of striated muscle. The association of modular protein-interacting domains enables ZASP to serve as an adaptor protein that recruits multiple proteins to a localized site of action in a specific subcellular region, for example α-actinin-2 and myotilin at the Z-discs of skeletal muscle (15, 36). Expression of mutant but not wild-type ZASP led to a loss of F-actin-cross-linking proteins, α-actinin-2, and myotilin, from the Z-discs and sarcoplasmic F-actin accumulation in skeletal muscle fibers. Time line studies in mice revealed that the loss of α-actinin-2 from the Z-discs preceded the sarcoplasmic accumulation of ZASP, α-actinin, and myotilin. Our results show that the wild-type and mutant sZM domains do not bind to α-actinin-2 and that the MFM mutations do not disrupt ZASP interaction with actin-cross-linking proteins. There remains a possibility that the MFM mutations in ZASP affect the binding affinity of α-actinin-2 to actin or another binding protein that is required for the integrity of the Z-disc structure in skeletal muscle.

The presence of multiple ZASP isoforms with unique biochemical roles and the isoform-specific effects of ZASP mutations suggest a molecular mechanism for zaspopathy, a late-onset autosomal dominant disease. ZASP haploinsufficiency is not likely, given that the measurement of ZASP by immunoblotting showed no clear difference in the expression of ZASP isoforms in skeletal muscle between patients and healthy controls (9) and given that heterozygous ZASP/Cypher knockout mice have normal muscle function and survival (14, 17, 31). Our results indicate that the MFM mutations in the sZM domain in actin-associated ZASP are deleterious to the Z-discs and F-actin organization in skeletal muscle fibers. This is likely not due to a direct effect of the interaction between the mutated sZM domain and F-actin because the mutations do not affect actin binding to the sZM domain and because the actin disruption is caused by the mutations in the ZASP-LΔex10 isoforms with two actin binding sites (the sZM domain and exon 8–11 junction) but not the isoforms that only contain the sZM domain. The isoform-specific phenotype indicates that ZASP-LΔex10 serves as a link between F-actin (via exon 8–11 junction site) and a binding partner other than F-actin for the sZM domain. Studies are ongoing to identify and characterize the binding partner(s) of the sZM domain that either are recruited specifically or bound more tightly to the mutated domain relative to the wild-type domain. Alternatively, the mutations may disrupt a physiological interaction of the wild-type domain that is important for the stability of the ZASP-actin-actinin complex at the Z-discs.

Mutations in at least five other Z-disc-associated proteins cause MFM with prominent disruption of skeletal muscle Z-discs (37). Although the cellular and molecular mechanisms underlying the Z-disc disruption in these myopathies have not yet been defined, some biological functions of the disease proteins are known. Myotilin, filamin C, and αβ-crystallin directly interact with F-actin, whereas desmin and BAG3 interact with the actin-associated proteins nebulin and CapZβ1, respectively (29, 38–41). Together with our findings of ZASP-actin interaction, it is likely that the molecular pathways leading to skeletal muscle Z-disc disruption are shared by these myopathies. Altered F-actin dynamics, either by direct or indirect associations of mutant proteins with actin filaments, may emerge as a unifying disease mechanism in this group of disorders.

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