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Laura T. Donlin, Christian Andresen, Steffen Just, et al.

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Smyd2 controls cytoplasmic lysine methylation of Hsp90 and myofilament organization

Laura T. Donlin,1 Christian Andresen,2 Steffen Just,3 Eugene Rudensky,1 Christopher T. Pappas,4 Martina Kruger,2 Erica Y. Jacobs,5 Andreas Unger,2 Anke Zieseniss,6 Marc-Werner Dobenecker,1 Tobias Voelkel,2 Brian T. Chait,5 Carol C. Gregorio,4 Wolfgang Rottbauer,3 Alexander Tarakhovsky,1,7,8 and Wolfgang A. Linke2,7

1Laboratory of Immune Cell Epigenetics and Signaling, The Rockefeller University, New York, New York 10065, USA; 2Department of Cardiovascular Physiology, Ruhr University, D-44780 Bochum, Germany; 3Department of Medicine II, University of Ulm, D-89081 Ulm, Germany; 4Department of Cellular and Molecular Medicine, The University of Arizona, Tucson, Arizona 85724, USA; 5Laboratory of Mass Spectrometry and Gasous Ion Chemistry, The Rockefeller University, New York, New York 10065, USA; 6Department of Cardiovascular Physiology, University Medicine Gottingen, D-37073 Gottingen, Germany

Protein lysine methylation is one of the most widespread post-translational modifications in the nuclei of eukaryotic cells. Methylated lysines on histones and nonhistone proteins promote the formation of protein complexes that control gene expression and DNA replication and repair. In the cytoplasm, however, the role of lysine methylation in protein complex formation is not well established. Here we report that the cytoplasmic protein chaperone Hsp90 is methylated by the lysine methyltransferase Smyd2 in various cell types. In muscle, Hsp90 methylation contributes to the formation of a protein complex containing Smyd2, Hsp90, and the sarcomeric protein titin. Deficiency in Smyd2 results in the loss of Hsp90 methylation, impaired titin stability, and altered muscle function. Collectively, our data reveal a cytoplasmic protein network that employs lysine methylation for the maintenance and function of skeletal muscle.

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Post-translational lysine methylation of histones plays a key role in the formation of chromatin-bound protein complexes that regulate a vast range of genome-related processes [Jenuwein and Allis 2001; Kouzarides 2007]. Lysine methylation has unique characteristics that contribute to the distinct functional properties of methyllysine-dependent protein networks. In comparison with other modifications, such as phosphorylation and acetylation, lysine methylation is considered a stable mark [Byvoet et al. 1972; Duerre and Lee 1974; Rice and Allis 2001]. In addition to its relative stability, the methyl mark can be found in one of three different forms. On histones, mono-, di-, or tri-methylation of individual lysines directs the association of functionally distinct chromatin proteins [Kouzarides 2007]. In addition to histones, nonhistone proteins in the nucleus, including p53, G9a, and NF-кB, have recently been demonstrated to incur lysine methylation (Chuikov et al. 2004; Huang et al. 2006; Sampaio et al. 2007; Ea and Baltimore 2009; Yang et al. 2009; Levy et al. 2010; Lu et al. 2010). Similar to histones, lysine methylation of nonhistone nuclear proteins contributes to formation of protein complexes (Sims and Reinberg 2008).

The role of lysine methylation in nuclear protein complex formation raises the question of whether methyllysine-dependent protein networks exist in the cytoplasm. The presence of the lysine methyltransferase (KMTase) Ezh2 in the cytoplasm and its role in cell surface receptor signaling have implicated a potential role for lysine methylation in the cytoplasm [Su et al. 2005]. However, the lack of information regarding an Ezh2 cytoplasmic substrate has precluded further understanding of how lysine methylation impacts cytoplasmic pathways.

In the current study, we demonstrate that the KMTase Smyd2, which localizes predominantly to the cytoplasm, methylates the chaperone Hsp90. We identify a specific site for Hsp90 lysine methylation and show that methylation of this site critically depends on Smyd2. We provide evidence that Smyd2 and methylated Hsp90 form a complex with the sarcomeric protein titin and that Smyd2 is required for normal function of skeletal muscle in vivo.

Results and Discussion

The Smyd2 KMTase localizes predominantly to the cytoplasm

Our search for putative cytoplasmic substrates of lysine methylation began with in silico and experimental screens for cytoplasmic KMTases. We reasoned that identification of cytoplasmic KMTases could lead to the discovery of cytoplasmic networks regulated by lysine methylation. The in silico screening of 40 KMTases showed that 11 contain a nuclear export signal [NES] and lack a nuclear localization signal [NLS], suggesting the potential for nonnuclear localization [Supplemental Table 1; Supplemental Fig. 1A]. These 11 KMTases were tagged with GFP and expressed in HEK 293 cells. We found that the enzymes Smyd2, Smyd4, and Set8/PR-Set7 predominantly localized to the cytoplasm [Supplemental Fig. 1A,B]. Furthermore, endogenous Smyd2 was also found in the cytoplasm of HEK 293 and C2C12 myoblast cells, as well as adult human skeletal muscle tissue [Supplemental Fig. 1C–E], which expresses high levels of Smyd2 in comparison with other tissues [Brown et al. 2006; Kawamura et al. 2008; Sun et al. 2008].

Smyd2 methylates the cytoplasmic chaperone Hsp90

The high abundance of Smyd2 in the cytoplasm and its ability to methylate the structurally distinct nonhistone...
nuclear proteins p53 and Rb (Huang et al. 2006; Saddic et al. 2010) rendered Smyd2 a favorable candidate to catalyze lysine methylation of cytosolic proteins. We therefore developed an approach to screen for cytosolic substrates of Smyd2. Cytosolic extracts were fractionated based on molecular weight, and each fraction was then incubated with recombinant Smyd2 and radiolabeled cofactor S-adenosylmethionine [SAM]. Using this method, we detected several Smyd2-methylated protein species that were then identified by MS/MS (Supplemental Fig. 2A,B). The most abundant Smyd2-methylated protein was found to be the chaperone Hsp90 (Fig. 1A, Supplemental Fig. 2B). Selectivity of Smyd2 for Hsp90 methylation was demonstrated by the inability of Smyd2 to methylate the Hsp70 chaperone or the endoplasmic reticular Hsp90 homolog gp96 (Fig. 1A; Supplemental Fig. 2C). Furthermore, the methyltransferases G9a and Set9, which have the capacity to methylate both histone and nonhistone proteins, failed to methylate Hsp90 in vitro (Supplemental Fig. 2D).

Using a panel of truncated Hsp90 proteins, as well as Hsp90 proteins in which individual lysine residues were replaced by alanine, we identified Lys 616 (K616) as the methylase and partial negative controls for methylated Hsp90, HEK 293 cells with or without exogenous Smyd2 expression were methyl Hsp90 (anti-Hsp90K616me1 antibody), Hsp90, and Smyd2. [shE] and nonsilencing shRNA [shN] analyzed by Western blot for lysates from U20S and C2C12 cells stably integrated with Smyd2-GFP and methylated Hsp90 to confirm that K616 represents a Smyd2 substrate, but also demonstrated that, in vivo, a single methyl group is covalently attached to K616, rendering it monomethyl K616 (K616me1) [Supplemental Fig. 3C]. K616 and the surrounding α-helix motif are highly conserved among Hsp90 homologs from mammals to yeast [Supplemental Fig. 3D]. Using a custom antibody specific for monomethyl K616 (anti-Hsp90K616me1) [Supplemental Fig. 3E], endogenous Hsp90K616me1 was detected in various cell lines and ex vivo isolated cells [Fig. 1C,D]. In particular, ex vivo isolated muscle tissues, which express the highest level of Smyd2 mRNA [Brown et al. 2006; Kawamura et al. 2008; Sun et al. 2008] and protein [Fig. 1D], also contained the highest levels of endogenous Hsp90K616me1 [Fig. 1D]. Multiple lines of evidence support an essential role for Smyd2 in the generation of Hsp90K616me1. First, methylation of Hsp90 was detected with the methyl Hsp90 antibody upon ectopic expression of Smyd2, but not the four other Smyd2 family members [Supplemental Fig. 3F]. Second, Hsp90K616me1 was undetectable in cells rendered Smyd2-deficient by expression of Smyd2-specific shRNAs [Fig. 1C].

**Figure 1.** Smyd2 methylates the Hsp90 chaperone. (A) Methyltransferase assay containing purified Hsp70, Hsp90, or histones and the radiolabeled cofactor SAM performed in the presence or absence of Smyd2. (B) Methyltransferase assays containing Smyd2 and C-terminal Hsp90 lysine-to-alanine [K–A] point mutants. Wild-type (Wt) and a dimerization domain-deleted (ΔDIM) Hsp90 were used as positive and negative controls, respectively. (His-Flx-Flag-tagged Hsp90 mutants; (e) endogenous copurifying Hsp90. (C) Whole-cell lysates from U20S and C2C12 cells stably integrated with Smyd2-specific shRNAs [sh1 and sh2] or two control vectors [empty vector [shE] and nonsilencing shRNA [shN]] analyzed by Western blot for methyl Hsp90 (anti-Hsp90K616me1 antibody), Hsp90, and Smyd2. HEK 293 cells with or without exogenous Smyd2 expression were used as positive and partial negative controls for methylated Hsp90, respectively. (D) Mouse tissue lysates analyzed by Western blot for methyl Hsp90, Hsp90, and Smyd2. [Br] Brain; [Thy] thymus; [Ht] heart; [Lu] lung; [Kid] kidney; [Spl] spleen; [SkM] skeletal muscle; [Liv] liver.

**Smyd2 and methylated Hsp90 colocalize on myofibrils**

The abundance of Smyd2 and Hsp90K616me1 in striated muscle suggests that methylation of Hsp90 by Smyd2 may have a function specific to myocytes. A GFP-tagged Smyd2 protein expressed in primary chick skeletal myocytes demonstrated a highly ordered striated arrangement, suggestive of myofibril binding [Fig. 2A]. Specifically, Smyd2 assembled as tightly spaced double bands flanking α-actinin striations that mark the sarcomeric Z-disks (Fig. 2A). Endogenous Smyd2 was also found in a similar pattern in human diaphragm muscle (Supplemental Fig. 4A). Incubation of purified human myofibrils with recombinant Smyd2-GFP protein, but not GFP alone, resulted in striations adjacent to Z-disks [Fig. 2B]. These data implicated direct binding between Smyd2 and myofibrils at the sarcomeric Z-disk, while other I-band components, such as actin, remain at a fixed distance (Linke et al. 1999; Cazorla et al. 2000). Upon myofibril stretching, we observed an increase in the distance between the Z-disk and Smyd2 striations [Fig. 3A]. For any given magnitude of stretching, this distance precisely matched that between the Z-disk and the N2A domain of titin, implying that Smyd2 interacts with this titin domain [Fig. 3A]. In support of an interaction, titin N2A striations colocalized with Smyd2-GFP [Supplemen-
tal Fig. 4B] and endogenous Smyd2 striations [Fig. 3B] in skeletal myocytes. A direct interaction between Smyd2 and the titin N2A domain was confirmed by yeast two-hybrid and GST pull-down assays, mapping the Smyd2-binding site to a unique sequence encoded by human titin exon 104 (Supplemental Fig. 4C,D). Despite the observed interaction, in vitro methyltransferase assays revealed no methylation of the titin N2A domain by Smyd2 (Supplemental Fig. 4E).

Hsp90, Smyd2, and titin form a protein complex dependent on Hsp90 methylation

Methylated Hsp90 colocalized with the same titin domain where Smyd2 binds [Fig. 3C], suggesting the possibility that Smyd2, methylated Hsp90, and the titin N2A domain are found together within a complex. Supporting this hypothesis, we found near-stoichiometric quantities of Hsp90 associated with overexpressed Smyd2 [Fig. 4A], as well as an interaction between endogenously expressed Smyd2 and Hsp90 [Fig. 4B], in C2C12 myoblasts. Furthermore, an exogenously expressed titin N2A domain associated with the Smyd2–Hsp90 complex in C2C12 cells. This association was identified under conditions in which Hsp90 or Smyd2 was immunoprecipitated, demonstrating the existence of complexes containing all three proteins [Fig. 4C; Supplemental Fig. 4F, respectively]. The association between Hsp90 and the titin N2A domain existed only in the presence of Smyd2 [Fig. 4C], demonstrating that Smyd2 is required for the Hsp90–titin interaction. In the absence of Smyd2-mediated methylation, the Smyd2–Hsp90 interaction was maintained, while titin binding to Hsp90 complexes was greatly reduced (Fig. 4C). These data suggest that Smyd2-mediated methylation of the chaperone Hsp90 promotes formation of a complex containing titin.

Smyd2-mediated Hsp90 methylation impacts titin protein stability

Hsp90 has been shown to be critical in myofibril organization and function [Barral et al. 2002; Du et al. 2008; Hawkins et al. 2008; Codina et al. 2010]. Although much of the evidence is related to its association with myosin, Hsp90 has also been implicated in titin function [Hawkins et al. 2008; Codina et al. 2010]. Having observed that methylation of K616 is necessary for the interaction of Hsp90 with titin, we noted that this lysine lies within a putative chaperone–substrate interface [Harris et al. 2004; Fang et al. 2006]. If titin represents a substrate of the chaperone Hsp90, one would predict that the interaction between methylated Hsp90 and titin would influence titin protein stability. C2C12 myoblasts expressing constant levels of titin N2A and increasing concentrations of Smyd2 cDNA showed increased methylation of endogenous Hsp90 and increasing concentrations of titin N2A proteins [Fig. 4C; Supplemental Fig. 4F, respectively]. The association between Hsp90 and the titin N2A domain existed only in the presence of Smyd2 [Fig. 4C], demonstrating that Smyd2 is required for the Hsp90–titin interaction. In the absence of Smyd2-mediated methylation, the Smyd2–Hsp90 interaction was maintained, while titin binding to Hsp90 complexes was greatly reduced (Fig. 4C). These data suggest that Smyd2-mediated methylation of the chaperone Hsp90 promotes formation of a complex containing titin.

Methylated Hsp90 stabilizes myofilaments

Figure 2. Smyd2 and methylated Hsp90 bind to muscle myofibrils. (A) Representative fluorescence images of embryonic chick skeletal myocytes expressing Smyd2-GFP. Arrows indicate a doublet of bands. Z-disks were visualized with an anti-α-actinin antibody. Bar, 10 μm. (B) Immunofluorescence images of isolated and stretched human cardiomyofibrils incubated with recombinant Smyd2-GFP and stained for α-actinin. Bar, 5 μm. (C) Immunofluorescence images of chick skeletal myocytes with or without Smyd2-GFP expression stained for methyl Hsp90 and α-actinin. Bar, 10 μm.

Figure 3. Colocalization and stretch-induced movement of Smyd2 with the titin N2A domain. (A) Distance from the Z-disk center for both Smyd2-GFP and the titin N2A domain (detected with a titin N2A antibody) in isolated human myofibrils stretched to increasing sarcomere lengths. (Open triangle and gray line) Smyd2-GFP; (solid square and gray line) Titin N2A; (F with arrow above) direction of stretch force. Bars, 3 μm. (B) Immunofluorescence images of chick skeletal myocytes stained for endogenous Smyd2, the titin N2A domain, and α-actinin. Bar, 10 μm. (C) Immunofluorescence images of human diaphragm muscle stained for endogenous methylated Hsp90 and the titin N2A domain (9D10 antibody). Bar, 5 μm.
protein [Fig. 4D]. This finding pointed to a positive role for Smyd2 in the regulation of titin stability. Expression of an enzymatically inactive Smyd2 resulted in decreased concentrations of titin N2A protein [Fig. 4D], indicating compromised titin protein stability. These data suggest that the enzymatic capacity of Smyd2 to methylate Hsp90 impacts titin protein stability. In support of this concept, we found that reduced Smyd2 expression negatively affected the structural integrity of the titin N2A domain. Treatment of myocytes with the Smyd2-specific siRNA resulted in a diminished striated staining pattern for the titin N2A domain [59% of cells with Smyd2 siRNA and 31% of cells with control siRNAs; n = 100, P = 6 × 10−4], while α-actinin striations appeared normal [Fig. 4E]. Reduced Smyd2 expression did not affect all regions of the giant titin filament (>1 μm) or induce titin protein degradation, as demonstrated by the wild-type titin pattern within the sarcomeric M-band region [Fig. 4E].

Smyd2 promotes stabilization of the sarcomeric I-band region

Our findings pointed to a potentially important role for Smyd2 in the regulation of muscle structure and function. Indeed, we found that reduced Smyd2 and methylated Hsp90 levels in zebrafish associated with skeletal and cardiac muscle defects [Fig. 5A]. Zebrafish with reduced Smyd2 expression displayed severely impaired mobility and contracted tails [Fig. 5B, C]. The impact on cardiac muscle performance was less severe, with reduced heart rates and fractional shortening decreased by ~50% [data not shown]. In an independent study on mouse cardiac development, Smyd2 was shown to be dispensable, suggesting a compensatory mechanism for the Smyd2 gene in higher-vertebrate heart development [Diehl et al. 2010]. The zebrafish phenotypes were observed upon attenuation of both Smyd2 alleles together (smyd2a and smyd2b) or with reduced smyd2a expression alone [Fig. 5B, C]. smyd2a expression was disrupted by two distinct nonoverlapping morpholino oligomers with the same phenotypic outcomes, but with variable penetrance (92.8% of n = 153 and 60% of n = 98). Histological analysis of zebrafish with reduced Smyd2 expression levels revealed disrupted skeletal muscle tissue [Supplemental Fig. 5A]. Electron micrographs demonstrated a localized and consistent disorganization of sarcomeric structures in the Z-disk and I-band regions [Fig. 5D; Supplemental Fig. 5B]. In contrast, the M-band regions maintained a normal ordered alignment in the absence of Smyd2 [Fig. 5D, Supplemental Fig. 5B]. The disordered I-bands and Z-disks in Smyd2-deficient zebrafish indicate that Smyd2 stabilizes the precise sarcomeric region where the Smyd2–Hsp90 complex is found.

In summary, we identified the cytoplasmic chaperone Hsp90 as a previously unknown substrate for the methyltransferase Smyd2. In muscle, we found that Smyd2-mediated methylation of Hsp90 regulated the formation of a complex with the sarcomeric protein titin. In this context, the formation of complexes around lysine methylation at the titin filament could be viewed as functionally analogous to the formation of complexes around lysine methylation at the chromatin fiber. In vertebrates, several Smyd family KMTases are expressed most highly in muscle cells (Gottlieb et al. 2002; Fuji et al. 2003; Tan et al. 2006; Kamamura et al. 2008; Sun et al. 2008; Thompson and Travers 2008). Thus, it is attractive to speculate that the Smyd family of KMTases will prove useful for understanding the role of these enzymes in the regulation of muscle structure and function.

Materials and methods

Methyltransferase assays

Reactions were performed in methyltransferase buffer (50 mM Tris at pH 8, 5 mM MgCl₂, 4 mM DTT) plus ~10 μM [3H]-adenosylmethionine (CE Healthcare or Perkin Elmer) and incubated for 45 min at 30°C. Protein gels
Methylated Hsp90 stabilizes myofilaments

Figure 5. Smyd2 regulates skeletal and cardiac muscle development and sarcomeric I-band structures in zebrafish. (A) Zebrafish lysates from control and Smyd2a/b morpholino-treated animals analyzed by Western blot for Smyd2, methyl Hsp90, and Hsp90. (B,C) Zebrafish injected with control or smyd2 antisense morpholino oligonucleotides. smyd2a and smyd2b were targeted in B, and only smyd2a was targeted in C. Images represent fish at 72 h post-fertilization. (D) Transmission electron microscopy of zebrafish skeletal muscle. (Z) Z-disk, (M) M-band. Bars: main panels, 1 μm, insets, 0.1 μm. Graph represents a blind scoring for the structural integrity of I-band/Z-disk and M-band regions. The percentage of regions with characteristic sarcomeric structure is plotted, with the error bars representing standard error. I-band/Z-disk: n = 38, P = 0.0002; M-band: n = 42, P = 0.3. (****) Statistical significance.

were stained with Coomassie, incubated with liquid EN’HANCE [Perkin-Elmer], incubated in PEG solution (10% polyethylene glycol 8000, 7% methanol, 7% acetic acid, 1% glycerol), dried, and exposed to Biomax XR film [Kodak] for 2–10 d at ~80°C.

Antibodies

The antibodies used were Smyd2 [Sigma], Flag M2 [Sigma], Hsp90 [Stressgen], Hsp90β [Stressman], and HA.11 [Covance]. The immunofluorescence antibodies used were α-actinin [Sigma], GFP [Abcam], Smyd2 [Eurogentec and Sigma], N2A-titin [Eurogentec], M-band titin [T114, custom-made], and Cy3- or Cy5-conjugated IgGs [Rockland].

Methyl Hsp90 antibody generation

A methyl Hsp90 peptide [NH2-NMERIMKme1AQALRDC] was synthesized by the Rockefeller University Proteomics Resource Center, injected into rats [Cocalico Biologicals], and affinity-purified [Open Biosystems].

Primary cultures of chick skeletal and cardiac myocytes

Breast muscle or heart from chicken embryos was minced and trypsinized, and nonadherent cells were plated on Matrigel-coated (BD) coverslips in growth medium (12.5% horse serum [HyClone], 12.5% chicken embryo extract, 5 mM glutamine in MEM [Cellgro]). After 2 d, growth medium was replaced by differentiation medium (12.5% horse serum, 2% chicken embryo extract) [Gregorio and Fowler 1995; Nawrotzki et al. 1995]. Transfections were performed using Escott IV [Sigma] or Effectene [Qiagen].

Mechanical manipulation and fluorescence staining of isolated myofibrils

Myofibrils were isolated from human donor hearts, prepared for mechanical measurements as described [Neagoe et al. 2002], and incubated with recombinant Smyd2-GFP [1.5 μg/μL buffer]. Distances from the Z-disk were measured under a Zeiss Axiosvert-135 microscope in epifluorescence mode (100× objective) using a CCD camera [Sony] and Imaged software.

Immunostaining of adult human diaphragm muscle

Muscle was fixed in 4% paraformaldehyde, permeabilized (0.5% Triton X-100), repeatedly washed, and incubated with primary and secondary antibodies. Samples were embedded in epoxy, thinly sectioned, and analyzed by confocal laser-scanning microscopy [Nikon, Eclipse Ti].

Smyd2 complex purification from myoblasts

C2C12 myoblast cells stably expressing His-Flag-tagged Smyd2 were lysed (0.1% NP-40, 20 mM HEPES, 100 mM NaCl, 1 mM MgCl2, 20 mM Na2MoO4, 2 mM Na3VO4, 30 mM NaF, protease inhibitor cocktail [Sigma], incubated with Flag beads [Sigma], washed with increasing amounts of salt [final wash containing 500 mM NaCl], and eluted with Flag peptides [Sigma].

Immunofluorescence microscopy of cultured myocytes

Myocytes were incubated in relaxing buffer [150 mM KCl, 5 mM MgCl2, 10 mM MOPS, pH 7.4, 1 mM EGTA, 4 mM ATP] and fixed in 2%–3% PFA. Images were captured with a Deltavision RT system [Applied Precision] and an inverted microscope [IX70, Olympus], a 100× objective, and a camera [CoolSNAP HQ, Photometrics] using SoftWoRx 3.5.1 software [Applied Precision].

siRNA Smyd2 silencing and analysis of titin N2A striations

Primary myocytes were transfected [Effectene, Qiagen] with 30 nM siRNAs [Ambion]. One-hundred cells were scored for distinguishable N2A striations, and a binomial distribution test was used for statistical analyses.

Zebrafish (Danio rerio) injection procedures and analysis

Morpholino-modified oligonucleotide sequences are listed in the Supplemental Material. Animals were scored positive for skeletal or cardiac muscle defects if they exhibited pericardial edema, elongated hearts, infold tract edema, reduced heart rate, curved tails, disorganized skeletal muscle fibers, and lack of myoseptum. Fish tail muscle was analyzed by transmission electron microscopy [Zeiss EM900] according to standard protocols. Three individuals blindly scored >30 sarcomeric I-band/Z-disc and M-band regions as either characteristic or disrupted in structure, with a Student’s two-tailed t-test used for statistical analyses.

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