Kdm3a lysine demethylase is an Hsp90 client required for cytoskeletal rearrangements during spermatogenesis

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ABSTRACT The lysine demethylase Kdm3a (Jhd2a, Jmjd1a) is required for male fertility, sex determination, and metabolic homeostasis through its nuclear role in chromatin remodeling. Many histone-modifying enzymes have additional nonhistone substrates, as well as nonenzymatic functions, contributing to the full spectrum of events underlying their biological roles. We present two Kdm3a mouse models that exhibit cytoplasmic defects that may account in part for the globozoospermia phenotype reported previously. Electron microscopy revealed abnormal acrosome and manchette and the absence of implantation fossa at the caudal end of the nucleus in mice without Kdm3a demethylase activity, which affected cytoplasmic structures required to elongate the sperm head. We describe an enzymatically active new Kdm3a isoform and show that subcellular distribution, protein levels, and lysine demethylation activity of Kdm3a depended on Hsp90. We show that Kdm3a localizes to cytoplasmic structures of maturing spermatids affected in Kdm3a mutant mice, which in turn display altered fractionation of β-actin and γ-tubulin. Kdm3a is therefore a multifunctional Hsp90 client protein that participates directly in the regulation of cytoskeletal components.

INTRODUCTION Normal development requires precision and sufficient plasticity to adapt to environmental and genetic changes. The recent discovery of the reversible nature of histone methylation has generated interest into two gene families encoding demethylase enzymes, as they play fundamental roles by mediating timely expression of developmental genes. This is illustrated by the disease phenotypes observed in animal models and human patients (Kooistra and Helin, 2012) associated with mutation in some of these genes. Jumonji domain (JmjC)-containing proteins form a large family of oxoglutarate-dependent dioxygenases capable of removing methyl groups from arginine and lysines of histones (Klose and Zhang, 2007). Knockdown of JmjC proteins gives rise to a wide range of phenotypes from embryonic lethality to no discernible abnormality (Takeuchi et al., 1995; Catchpole et al., 2011; Iwamori et al., 2011; Ishimura et al., 2012). Despite the fact that many histone demethylases bind to a large number of genes, their silencing or knockdown alters the expression of relatively few genes with rather modest changes in expression levels. This suggests that histone demethylases may not have a major role in the regulation of gene expression but rather may be fine modulators of chromatin epigenetic states (Kooistra and Helin, 2012). Although normal development can be...
potentially disrupted by the synergistic effect of small changes in the expression of functionally related genes, the modest effect on transcription raises the interesting possibility that many histone demethylases may alter development through nonhistone functions as well (Webby et al., 2009; Lu et al., 2010).

Kdm3a (Jhmd2a, Jmjd1a, KIAA0742, TSGA) is a jumonji-domain protein able to remove mono- and dimethyl groups of histone H3 Lys-9 (H3K9me1 and H3K9me2) (Yamane et al., 2006). Kdm3a knockdown leads to immotile round-headed spermatozoa (globozoospermia) and metabolic syndrome associated with obesity and diabetes (Okada et al., 2007; Inagaki et al., 2009; Tateishi et al., 2009; Liu et al., 2010) through the regulation of genes involved in histone replacement during spermatogenesis and regulation of gene expression during adipogenesis. It has recently been shown that lack of Kdm3a leads to male-to-female sex reversal at non-Mendelian ratios, through misregulation of the sex-determining gene sry (Kuroki et al., 2013). Kdm3a is highly expressed in testis and at lower levels in fat tissue and can be readily purified from HeLa cells (Tskudaka and Zhang, 2006). Kdm3a was shown to be a coactivator of estrogen and androgen receptors (Yamane et al., 2006) and sry (Kuroki et al., 2013); further studies showed that Kdm3a is also involved in many cellular processes, including progression through the cell cycle (Cho et al., 2011), embryonic (Loh et al., 2007) and adult (Ma et al., 2008) stem cell renewal, and differentiation of vascular smooth muscle (Lockman et al., 2007) and parietal endoderm (Herzog et al., 2012). Increased methylation of histone H3K9 in tests after Kdm3a knockdown has been found when spermatids are fractionated by stage (Liu et al., 2010) but not when whole testis extracts are analyzed (Okada et al., 2007), illustrating the stage-specific and modulatory nature of this demethylase. Nevertheless, global depletion of mono- and dimethylated histones is consistently observed following Kdm3a overexpression or in vitro incubation with purified Kdm3a protein (Yamane et al., 2006). Thus the function of Kdm3a as a histone lysine demethylase has been extensively demonstrated.

Kdm3a was shown to participate in the transcriptional control of transition proteins and proteamines leading to postmeiotic chromatin condensation defects (Okada et al., 2007; Liu et al., 2010). It is thus postulated that the lack of DNA condensation underlies the abnormal sperm head shaping in Kdm3a animal models (Okada et al., 2010). The formation and shaping of the mammalian sperm head is a complex process that involves sequential changes in the head of the nucleus forming the head-tail coupling apparatus (HTCA). At step 9, the nucleus starts to elongate, forming a dorsal and ventral surface in tight association with the acrosome and manchette microtubules forming the peripheral ring that delineates its intersection with the acroplaxome (see diagram in Figure 2A later in this article). These tubulin- and actin-based structures play major roles in shaping the spermatid head by providing a structural platform for clutching forces during nuclear elongation (Kierszenbaum et al., 2003). The most common causes of round-headed spermatozoa are often associated with acrosomal and manchette defects (Dam et al., 2007). The intrinsic complexity of sperm head shaping renders difficult the assignment of a single molecular event to the causation of globozoospermia. Interestingly, some endogenous Kdm3a protein is present in the cytoplasm during spermatogenesis (Okada et al., 2007), but the potential contribution of non–histone-related roles for Kdm3a during spermatogenesis has not been investigated.

In this paper, we show that Kdm3a mutant mice have cytoplasmic defects preceding histone replacement and chromatin compaction that significantly contribute to arrest spermatid elongation and produce rounded sperm heads as previously reported for these models (Okada et al., 2007; Liu et al., 2010). Electron microscopy shows that elongating spermatids in homozygous animals lacking Kdm3a catalytic domain (Kdm3aΔJC) have deformed acrosomes, incomplete manchettes, and detached centrosomes. Similar but milder features are observed in a second mouse model with a gene-trap insertion within Kdm3a (Kdm3aΔJ), which presents normal electro-dense (condensed) DNA but spermatid heads that are still rounded. We confirm the existence of a short Kdm3a isoform that is enzymatically active, expanding the complexity of Kdm3a regulation. This isoform is not interrupted by the gene-trap insertion, and although transcript levels are very low, this isoform may contribute toward the phenotype amelioration observed in the second model. We found Kdm3a to interact with the cellular chaperones: Hsp90, Cct/Tric, and Vcp. Multiple independent experimental designs addressed the specificity of Kdm3a interaction with Hsp90 and its requirement for Hsp90 chaperoning for its demethylase activity. Antibodies to an Hsp90 lysine residue known to be dynamically methylated (Abu-Farha et al., 2011; Donlin et al., 2012) indicate that this site is not modified by Kdm3a. Nevertheless, Hsp90 showed altered association with cytoplasmic components in somatic and germ cells of Kdm3a mutant mice, providing evidence that the cytoskeletal defects are a direct consequence of inactive Kdm3a. Our work provides molecular evidence for a previously unknown role of Kdm3a in the extensive cytoskeletal rearrangements required for spermatogenesis to proceed normally.

RESULTS

Kdm3a mouse models

To expand the functional studies of Kdm3a, we generated two mouse models. First, we rederived a previously published targeted mutant allele that lacks the Kdm3a catalytic domain JmJC (Kdm3aΔJC; Tateishi et al., 2009; Figure 1A). In previous studies, these mice provided evidence for the role of Kdm3a in the regulation of metabolic gene expression and resistance to obesity (Tateishi et al., 2009). homozygous males were shown to be infertile, with testes significantly smaller than those of wild-type littermates (Okada et al., 2007). Mice containing recombination Kdm3a alleles (ΔJC) were identified by PCR (Figure 1, A and B). Western blot analysis with a Kdm3a-specific monoclonal antibody directed to the N-terminal end of Kdm3a not contained within the deletion (Figure 1A, pAb N-Kdm3a) revealed the absence of full-length protein in homozygous cells and the appearance of low levels of deletion product in heterozygous and homozygous cells (Figure 1C). As shown previously, epididymis of Kdm3aΔJC/ΔJC animals shows a drastic decrease of spermatozoa and the few found have deformed heads (Figure 1D). Heterozygous animals were fertile (Table 1), but inspection of testis cross-sections revealed aberrant accumulation of large nuclei in the lumen of seminiferous tubules (Figure 1E and F).

The second mutant mouse allele contains a gene-trap insertion (Kdm3a57) within intron 5–6 of the Kdm3a locus (Figure 1G). Heterozygous animals derived from germ line–transmitting male
FIGURE 1: Two Kdm3a mouse models present arrested spermatogenesis with globozoospermia. (A) Diagram illustrates Cre-mediated deletion of exons 22–24 containing JmjC catalytic domain of Kdm3a in Kdm3aΔJC mice (Tateishi et al., 2009). (a), (b), and (c) indicate the position of primers used to genotype mutant mice. Color code: green oval, C6-type zinc finger (ZF); blue, LXXLL nuclear receptor binding; red, JmjC domain; black, exons surrounding and affected by deletion. pAb, polyclonal antibody. (B) Genomic PCR identifying ΔJC deletion mutants. (C) Immunoblot of total MEF extracts with Kdm3a antibody (N-terminal) shows absence of full-length Kdm3a protein (FL, red arrow) in Kdm3aΔJC/ΔJC. A shorter Kdm3a protein product result of Cre-induced deletion is indicated (ΔJC, black arrow). A nonspecific band is indicated (*). (D) DAPI staining of Kdm3aΔJC/ΔJC epididymis reveals few mature sperm with rounded nuclei. (E) DAPI staining of testis cross-sections shows some abnormal tubules in Kdm3aΔJC/+ (boxed region). Kdm3aΔJC/+ show arrested spermatogenesis. (F) Higher magnification of indicated regions from (E). es, elongated spermatids; arn, accumulation of...
chimeras were backcrossed onto C57BL6J females. Reverse transcriptase followed by PCR and Western blot analysis revealed that Kdm3a<sup>GT/GT</sup> mice lack Kdm3a full-length transcript (Figure 1H) and protein (Figure 1I). Kdm3a<sup>GT/GT</sup> males are infertile, whereas heterozygous animals produce offspring (Table 1). Kdm3a<sup>GT/GT</sup> tests are also significantly smaller than those of wild-type and heterozygous littermates (Figure 1J). Cross-sections of Kdm3a<sup>ΔJC/ΔJC</sup> and gene-trap (GT) mice. Each genotype is present in the expected Mendelian ratios.

**TABLE 1**: Heterozygous Kdm3a animals are fertile.

<table>
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<tr>
<th>Mouse model</th>
<th>Genotype (%)&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Total born</th>
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<tr>
<td>ΔJC</td>
<td>23</td>
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<td>GT</td>
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<sup>a</sup>Percentage of genotypes in offspring from heterozygous crosses of knockout (ΔJC) and gene-trap (GT) mice. Each genotype is present in the expected Mendelian ratios.

Cloning of a new murine Kdm3a isoform

There are several predicted isoforms for mouse and human Kdm3a (Ensembl). The milder severity of Kdm3a gene-trap mice prompted us to consider the existence of an alternative isoform not affected by the gene-trap insertion that could partially compensate for the lack of the full-length protein. We designed primers following a prediction from Ensembl in which a transcript with a unique exon contained within intron 11–12 would encode for a short Kdm3a isoform (Figure 4A). Using mRNA from Kdm3a<sup>ΔJC/ΔJC</sup> tests, we cloned a properly spliced transcript containing the three major Kdm3a domains and an alternative exon not found in the full-length transcript (Figure 4B, unique exon). Quantitative RT-PCR shows that transcript levels of this isoform, named isoform-2 (i2) is unaltered in adult testis from Kdm3a<sup>ΔJC/ΔJC</sup> mice (Figure 4C), indicating that the gene-trap insertion does not affect its expression in this tissue. Kdm3a-i2 is therefore likely transcribed from a promoter downstream from the gene-trap insertion. However, RNA expression of Kdm3a-i2 is much lower than full-length Kdm3a relative to gliceraldehyde 3-phosphate dehydrogenase (GAPDH; Figure 4C). For testing whether Kdm3a-i2 encodes an enzymatically active histone demethylase, Kdm3a-i2 was cloned in-frame with a green fluorescent protein (GFP) tag and transfected into human retinal pigmented epithelium cell line RPE1 (Supplemental Figure S1A). The overexpressed protein localized to
FIGURE 2: Ultrastructural defects of Kdm3aΔJC/ΔJC precede and persist after chromatin condensation. (A) The diagram illustrates cellular structures observed by electron micrographs. (B) Representative electron micrograph of a wild-type (WT) spermatid in early step 9. The acrosome (ac) has flattened and spreads symmetrically over the nuclear envelope. Microtubules of the manchette (dotted line) emanate from the perinuclear ring (pr) parallel to the nuclear surface. The caudal end of the nucleus is indented by the implantation fossa (if), with the mother centriole closely apposed (mc). (C) Step 9 spermatid in homozygous testis shows absence of ventral acrosome (va) surface (red arrow) and microtubules of the manchette (mm, dotted line) extending beyond the perinuclear ring (pr). Caudal end of nucleus remains round without traces of implantation fossa (if) in this or other planes of this spermatid section. Electrodense portions in nucleus represent condensing chromatin (cc). (D) Low magnification of homozygous sections illustrating the extent of abnormal spermatids. Golgi has moved to caudal end (go; spermatid 1). One-sided acrosome without ventral acrosome (va; 2). Strongly condensed chromatin (cc; 3). Disrupted manchette (4). (E) Spermatid 4 shown in (C) presents disrupted perinuclear ring (pr) with only a few manchette microtubules (mm). Several cc domains are observed. (F) Condensed spermatid from homozygous animals displays dark nucleus, sign of chromatin compaction and if impacted by HTCA. Scale bars: 2 μm.
the nucleus and cytoplasm (Figures 4C and S1B) regardless of whether the tag was positioned at the N- or C-terminal end of the protein (unpublished data). Decreased immunostaining of transfected cells with antibodies to lysine-methylated histones indicates that this newly cloned Kdm3a isoform is also an active histone lysine demethylase (Figure 4, D and E).

In summary, we show that Kdm3a encodes at least two enzymatically active isoforms. This newly identified protein could provide partial rescue, explaining the decreased severity of Kdm3a<sup>ΔJC/ΔJmc</sup> mice.

**Identification of Kdm3a-interacting proteins in testis**

Originally, Kdm3a was isolated in a two-hybrid screen using different domains of Hsp90 as baits. Hsp90 chaperones, one of the most abundant cellular proteins, play multiple roles assisting client proteins that must undergo large structural changes for biological activity (Kim et al., 2013). We used different domains of Hsp90aa1 (Hsp90α) and Hsp90ab1 (Hsp90β) and found that the catalytic domain of Kdm3a interacts with the middle region of Hsp90ab1. The N-terminal and C-terminal dimerization domains of Hsp90aa1 failed to interact with Kdm3a under these experimental conditions (Figure 5A). Kdm3a interaction with Hsp90 was further validated through glutathione S-transferase (GST) pull downs, which showed that both the catalytic JmjC domain (GST-JmjC) and a C-terminal truncation without the catalytic domain (GST-ΔJmjC) interact with Hsp90, albeit with different efficiency as determined with pan-Hsp90 antibodies (Figure 5B).

To investigate the interaction profile of endogenous Kdm3a, we immunopurified testis extracts with antibodies to Kdm3a; this was followed by mass spectrometry (MS). We used extracts from wild-type and Kdm3a<sup>ΔJC/ΔJmc</sup> animals. Three independent pull downs for each genotype from mice age 1.5–6 mo were resolved in polyacrylamide gels, and the indicated regions were cut and analyzed by MS (Figure 5C). In agreement with two-hybrid studies, we found multiple cellular chaperones from the Hsp90 family; we also found Cct chaperonins in Kdm3a pull downs absent in control samples (Figure 5D). There is also abundant representation of the chaperone valosin-containing protein Vcp and the RNA-binding protein Fxr1. Kdm3a interaction with Hsp90 was further validated by immunoblotting with antibodies specific to Hsp90a1 and Hsp90ab1. Pull-down extracts from wild-type and heterozygous mice specifically contain Hsp90ab1 and Hsp90a1, indicating that Kdm3a interacts with both chaperones (Figure 5E). Interestingly, the truncated Kdm3a protein of Kdm3a<sup>ΔJC/ΔJmc</sup> mice is bound and immunopurified by an antibody directed to the N-terminal region of Kdm3a (Figure 5E and F). As determined by the relative abundance of Hsp90 to immunopurified Kdm3a proteins, the truncated form of Kdm3a binds Hsp90 chaperones more efficiently or in greater proportion than the full-length Kdm3a protein. Importantly, the catalytic domain of Kdm3a found in the two-hybrid screen to interact with Hsp90ab1 (Figure 5A) is deleted in Kdm3a<sup>ΔJC/ΔJmc</sup> mice (Figure 1A), but, as seen with the truncated GST-ΔJmjC construct, the C-terminal end of Kdm3a contributes to this interaction (Figure 5B). This observation suggests that multiple residues are responsible for Kdm3a interaction with Hsp90; alternatively, as Kdm3a may be rendered unstable through the deletion, it may have increased affinity for Hsp90.

In summary, these results show that Kdm3a interacts specifically with Hsp90 chaperones in vivo.

**Substrate or client?**

Kdm3a interaction and regulation of cytosolic chaperones would be consistent with the ultrastructural defects observed in Kdm3a mouse models. The chaperonin containing TCP1 (t-complex poly peptide 1/Cct/TriC) forms two heterogeneous eight-membered complexes (CCT-1-8) (Dekker et al., 2011) that mediate the folding and assembly of a small subset of nascent polypeptides, preventing their aggregation within the crowded cytoplasmic environment (Kim et al., 2013) and playing a significant role in the cytoskeletal rearrangements required to produce mature spermatids (Willison et al., 1989; Sternlicht et al., 1993; Souès et al., 2003). Cct chaperonins are essential for the de novo folding of actin and tubulin (Llorca et al., 2000), which form part of the acroplaxome and microtubular manchette affected in Kdm3a mouse models. Likewise, a critical requirement for Hsp90 in the extensive microtubular rearrangements during spermatogenesis has been documented (Yue et al., 1999), as has Hsp90 functional interplay with CCT chaperonins (Kim et al., 2013). Moreover the chaperone functions of Hsp90 are regulated by multiple posttranslational modifications (Li and Buchner, 2013), including lysine methylation by SmyD2 (Abu-Farha et al., 2011). This modification is in turn important for myocyte organization (Donlin et al., 2012). Misregulation of the posttranslational state of these cellular chaperones could thus contribute to the observed cytoplasmic defects of Kdm3a mutant mice. Conversely, Hsp90 could regulate Kdm3a functions instead, and the structural defects observed could be a consequence of the direct participation of Kdm3a on cytoplasmic components. For example, Hsp90 was found to modulate the stability of Kdm4b histone demethylase (Ipenberg et al., 2013) and to enhance the histone methyltransferase activity of SmyD2 (Abu-Farha et al., 2008).

We first investigated the methylation state of Hsp90 in the absence of Kdm3a demethylase activity through the use of tailored antibodies to methylated K616 (Hsp90 meK616) of Hsp90 (Donlin et al., 2012). Individual pull downs of Hsp90aa1 and Hsp90ab1 indicate that both Hsp90 chaperones are methylated in vivo (Figure 6A). Neither total extracts (Figure 6B) nor Hsp90 pull downs (Figure 6C) from Kdm3a<sup>ΔJC/ΔJmc</sup> testis show significant changes in the methylation levels of Hsp90 as detected with meK616 antibody, indicating that Kdm3a does not regulate the methylation state of this lysine residue. It has been shown, however, that Hsp90 has multiple methylation sites (Abu-Farha et al., 2011), and it thus remains possible that other methylated lysines of Hsp90 could be regulated by Kdm3a.

We then investigated whether Kdm3a requires Hsp90 for its demethylase activity, using histone methylation as a readout. Red fluorescent protein (RFP)- or GFP-tagged full-length or i2 Kdm3a (Supplemental Figure 1A) were transfected in parallel with their tag control. Six hours later, the cells were treated with the Hsp90 inhibitor 17AAG. The dimethylated state of histone H3 Lys-9 (H3K9me2) was assessed 48 h after transfection. Both tagged isoforms display nuclear and cytoplasmic distribution, albeit Kdm3a-FL nuclear presence is very strong (Supplemental Figure 1B). We observed a marked decrease in the intensity of tagged-Kdm3a upon treatment with Hsp90 inhibitors and a shift in the subcellular distribution, suggesting Hsp90 may regulate Kdm3a protein stability and localization. To dissociate the loss of Kdm3a demethylase activity from these observations, we measured H3K9me2 levels in cells with high levels of tagged-Kdm3a only (Figure 6D). These results indicate that the enzymatic activity of both Kdm3a isoforms depends on Hsp90, as their demethylation capacity decreases when Hsp90 is inhibited. The effect of Hsp90 on Kdm3a levels was confirmed by treating RPE1 cells with 17AAG followed by immunodetection of endogenous KDM3A protein (Figure 6E). The change in subcellular distribution was investigated by determining the ratio of nuclear to cytoplasmic Kdm3a distribution was investigated by determining the ratio of nuclear to cytoplasmic Kdm3a.
**FIGURE 3:** Kdm3a gene-trap model presents milder structural defects. (A) Representative electron micrograph of a wild-type (WT) spermatid in early step 9. (B) Homozygous animals have ventral (va) and dorsal (da) acrosome surfaces, but manchette descends asymmetrically (arrows), and nuclear membrane has folds (fld). (C) The mother centriole (ce) is engaged in implantation fossa (if) in WT condensed spermatids. (D) Condensed spermatids from homozygous animals have detached mother centriole (ce) without clear implantation fossa (if) and detached electrondense material (em). (E) Low magnification of a homozygous-GT tubule illustrates the extent and frequency of condensed spermatids. (F) Epididymal spermatozoa from heterozygous animals stained with septin 4 (annulus and acrosomal marker, green) show acrosome expanding over a normal sperm head. Homozygous sperm shows cube-like head with horizontal acrosomes. Scale bars: A–E, 2 μm; F, 10 μm.

**FIGURE 4:** Kdm3a encodes two protein isoforms. (A) Intron-exon diagram of Kdm3a. Short isoform i2 was cloned following ENSEMBL prediction (ID numbers between parentheses). First and unique Kdm3a-i2 exon is contained within intron 11. (B) Diagram illustrates the amino acid position of Kdm3a protein domains in full-length (FL) and isoform 2 (i2). Green line, i2-exclusive exon; amino acid sequence shown in bold. pAb indicates the relative position of amino terminal Kdm3a single polyclonal antibody used along this study. (C) qRT-PCR from testis and MEF RNA show a decrease of Kdm3a-FL and unaltered transcript levels of i2 in Kdm3aGT/GT testis. A slight reduction of Kdm3a-i2 is observed in Kdm3aGT/GT primary MEFs. (D) RPE1 cells transfected with GFP tag fused upstream of Kdm3a-i2. Monoclonal antibodies to mono- or dimethylated Lys-9 of histone H3 were used to determine the state of histone methylation (red) in transfected cells (green). Arrows point to transfected cells. (E) Quantitation of (D). n = number of cells counted for each condition. *p < 0.05, Fisher’s exact test.
FIGURE 5: Kdm3a interacts with Hsp90 and other cellular chaperones in vitro and in vivo. (A) Positive clones identified from a large two-hybrid screen using Hsp90 as baits were purified, retransformed into yeast, and mated to reassess interactions. The catalytic domain of Kdm3a can rescue amino acid deficiency of the parental yeast strain only when Hsp90ab1 is the bait. Diagram illustrates JmjC region contained in two-hybrid and GST constructs. (B) Bacterially expressed C-terminal end of Kdm3a-GST: GST-JmjC (aa 1136–1323), GST-JmjCΔ (aa 1283–1323), and GST vectors were incubated with HeLa cell extracts. Proteins bound to glutathione beads were resolved and blotted with the indicated antibodies to determine interactions of GST constructs with Hsp90. (C) TrueBlue stain of whole adult testis extracts immunopurified with Kdm3a or an irrelevant antibody used as control (Ir Ab). The gel section indicated was cut for mass spectrometry (MS). (D) Table shows the number of unique peptides identified with >99% confidence for three independent immunopurifications resolved as shown in (C) for each genotype. Only proteins shared by the six runs and not found in control samples are included. (E) An antibody directed to the N-terminal end of Kdm3a pulls down endogenous Hsp90 from testis. Note that the in-frame deletion Kdm3a protein encoded by the floxed Kdm3a allele still binds Hsp90aa1 and Hsp90ab1 strongly. (F) Total extracts from adult testis immunoblotted with Kdm3a show absence of full-length (FL) and low levels of truncated (ΔJC) Kdm3a protein in Kdm3aΔJC/ΔJC animals. Kdm3a-FL (red arrow) in wild-type (WT) and heterozygous (Het) but not control pool is indicated. 1–3 identify an animal for each genotype.
cytoplasmic fluorescence intensity of Kdm3a-transfected cells grown in solvent control or in the presence of 17 AAG (Figure S1C). These measurements confirm a significant shift of RFP-Kdm3a-i2 but not RFP-Kdm3a-FL toward the cytoplasm upon Hsp90 inhibition (Figure 6F and G).

These results uncover a complex dependence of Kdm3a on Hsp90 to fulfill its biological functions. Interestingly, Hsp90 did not significantly affect the nuclear-cytoplasmic distribution of the full-length Kdm3a isoform but altered the distribution of Kdm3a-i2. This is perhaps because the short isoform (i2) does not contain the N-terminal region encoding a previously described nuclear localization signal (Brachule et al., 2013). The dependence of both isoforms on Hsp90 for their enzymatic activity therefore suggests a specific requirement of the Hsp90 chaperone machinery for the demethylation reaction. Moreover, the absence of detectable changes in the methylation state of Hsp90 in Kdm3a mutant mice suggests the cytoskeletal defects uncovered in this study may reflect a direct role of Kdm3a in the cytoplasm rather than these being a consequence of Kdm3a-mediated dysregulation of the chaperone system in Kdm3a mutant mice.

Cytoplasmic localization of Kdm3a

Previous publications have observed cytoplasmic distribution of Kdm3a by means of immunofluorescence of testis sections (Okada et al., 2007) and somatic cells (Okada et al., 2007; Sar et al., 2009; Yang et al., 2009; Yamada et al., 2012). We extended these findings using a fractionation kit (Qproteome) that allows the sequential isolation of soluble and organelle-bound proteins from cytoplasm and nucleus, respectively. Subcellular fractionation of mouse embryonic fibroblasts (MEFs) followed by immunoblotting shows nuclear and cytoplasmic (Figure 7A, 0.25% FCS) distribution of Kdm3a. There is also a marked shift though the nuclear signal of the tagged protein is stronger (Figures 7A, 10% FCS). Finally, the use of an antibody directed to the C-terminal end of Kdm3a that does not recognize 10% FCS). The endogenous Kdm3a full-length protein is nuclear and cytoplasmic in exponentially growing cultures (Figure 7A, 10% FCS), as observed for transfected RFP-KDM3A-FL in exponentially growing RPE1 cells, although the nuclear signal of the tagged protein is stronger (Figures 6F and S1B). In resting cells, however, Kdm3a is almost exclusively cytoplasmic (Figure 7A, 0.25% FCS). There is also a marked shift between soluble and membrane-bound Kdm3a depending on the growth conditions (Figure 7A, compare lanes 1 and 2 of 0.25 and 10% FCS). Finally, the use of an antibody directed to the C-terminal end of Kdm3a that does not recognize Kdm3a truncated protein (Figure 7B) shows Kdm3a presence in the cytoplasm with strong enrichment in the acrosome of wild-type but not Kdm3a mature spermatids (Figure 7C).

These results indicate that Kdm3a distribution within cells is dynamic and influenced by growth conditions. These findings are consistent with the hypothesis that Kdm3a plays a direct role in the cytoplasm of maturing spermatids.

Investigation of the role of Kdm3a in the cytoplasm

To begin to explore the cytoplasmic role of Kdm3a, we used three independent approaches, bearing in mind that Kdm3a could interact with but might not necessarily demethylate nonhistone substrates. We reasoned that, if Kdm3a requires Hsp90 for its biological functions, differential associations of these chaperones between controls and homozygous animals evidenced by partition along sucrose gradients or IP/MS would provide information on the state of the cytoskeleton and potential Kdm3a-interacting proteins in the cytoplasm. Likewise, if Kdm3a has nonhistone substrates in the cytoplasm, these would be evidenced as differentially methylated proteins between Kdm3a homozygous and control animals.

Actin and tubulins are among the most abundant components of the acrosome and manchette found in multiple polymerization states (Kierszenbaum et al., 2011). Their dependence on Cct and Hsp90 chaperones for maturation led us to explore their state in Kdm3a ΔJC/ΔJC mutant tests. Total tests extracts show comparable levels of Hsp90ab1 and Cct4 used as indicators of Hsp90 and chaperonin II complex, respectively (Figure 8A). Levels of β-actin and γ-tubulin are also seemingly unaltered (Figure 8A). Sucrose gradients of cytoplasmic extracts from tests show slightly altered fractionation of chaperonins and Hsp90 in Kdm3a ΔJC/ΔJC (Figure 8B). Hsp90ab1 migration along Kdm3a ΔJC/ΔJC gradients is more compact, while that of Cct3 is more widespread compared with wild-type gradients (Figures 8B and S2). Most striking is the differential partition of γ-tubulin and β-actin. In homoygous tests, γ-tubulin is found in early fractions, suggesting that a significant amount of the cellular pool of γ-tubulin dissociates easily or is not part of large polymers (Figures 8B and S2, red lines). This is illustrated by measuring the relative distribution of γ-tubulin along each gradient (Figure 8C). On the contrary, β-actin shows increased presence in late fractions of the homozygous samples, indicating that in these mice actin is part of larger complexes, perhaps aggregates (Figure 8B, blue line, and D). These gradients also confirm the presence of Kdm3a in the cytoplasm, as detected with the two polyclonal antibodies validated in this study (Figures 8B and S3A). The altered state of cytoskeletal components relative to chaperone complexes in Kdm3a homozygous animals also extends to somatic cells (Figure S4).

Immunopurification of total tests extracts with antibodies to methylated lysines revealed a single band differentially present in Kdm3a ΔJC/ΔJC samples that belongs to the actin family, β-actin–like protein 2 (k-actin or Actb2; Figure 8D). One of the identified peptides for this protein presents evidence of a potentially methylated lysine residue (Figure S5A). We then immunopurified Hsp90 chaperone complexes from age-matched wild-type, Kdm3a ΔJC/ΔJC, and Kdm3a ΔJC/ΔJC tests; this was followed by MS. Hsp90 interactions are usually transient; protein extracts were therefore incubated with antibodies for 1 h in the presence of sodium molybdate, thereby freezing the Hsp90 chaperone cycle and stabilizing interactions with client proteins (Taipale et al., 2012). This provides a snapshot of the state of Hsp90 complexes for each genotype. These experiments show largely overlapping hits for Hsp90 pull downs between the three Kdm3a genotypes absent in all the controls (WT, Kdm3a ΔJC/ΔJC, and Kdm3a ΔJC/ΔJC) performed with an irrelevant antibody (Supplemental Table S2). Actb2 appears to be found exclusively in Kdm3a ΔJC/ΔJC pull downs, but the previously identified methylated peptide was not found, and none of the remaining peptides found present evidence for methylation (Figure S5B). Total extracts immunoblotted with Actb2 antibodies show that the overrepresentation of Actb2 in Kdm3a ΔJC/ΔJC pull downs is not due to increased expression levels (Figure 8F). Cct chaperonins also seem enriched in Hsp90 pull downs from Kdm3a ΔJC/ΔJC extracts, particularly Cct7, which is found only in Kdm3a ΔJC/ΔJC samples (Table S3). These findings are consistent with the increased overlap between Hsp90 and Cct chaperonins observed in Kdm3a ΔJC/ΔJC gradients (Figure S2D) and the initial finding of Cct proteins in Kdm3a pull downs (Figure 5D). Furthermore, among all the methylated peptides identified in Hsp90 pull downs, 11 were found in Kdm3a ΔJC/ΔJC samples only (Table S3). From these, Cct2 (Ttc chaperonin 1 beta; Figure 5C) and the cytoplasmic dyneins Dc111 and Dyhc1 could directly contribute to the cytoplasmic defects observed in Kdm3a mutant tests.

These three independent approaches provide consistent evidence on the altered state of cellular chaperones and on the
FIGURE 6: Kdm3a is an Hsp90 client protein. (A) Total extracts from ΔJC Kdm3a testis were immunopurified with the indicated Hsp90 antibodies and immunoblotted with an antibody to methylated Lys-616 (Me K616) of Hsp90. TrueBlue staining shows the gel fraction used to confirm these pull downs by MS. (B) Total extracts from testis immunoblotted with the indicated antibodies show comparable levels of methylated Hsp90 as detected with Me K616 antibody. (C) The same unaltered trend in methylation is observed when antibody-purified Hsp90 pull downs are blotted. A seemingly
 FIGURE 7: Cytoplasmic distribution of endogenous Kdm3a. (A) Subcellular fractionation of serum-starved and exponentially growing MEFs immunoblotted with the indicated antibodies. Sequential disruption of cell and nuclear membranes gives rise to two cytoplasmic fractions: soluble (1) and membrane (2) and two nuclear fractions: soluble (3) and cytoskeletal (4). GAPDH and HP1 are controls for cytoplasmic and nuclear fractions, respectively. Red and black arrows indicate FL-Kdm3a and GAPDH proteins, respectively. (B) Total extracts from four wild-type (WT) and 6 ∆JC Kdm3a mice immunoblotted with pAb antibody directed to the C-terminal region of Kdm3a (see also Figures 6B and S2A). The immunizing peptide of this commercially available antibody is likely contained within ∆JC deletion, as no band is observed in any of the three Kdm3a∆JC/∆JC mice analyzed. GAPDH is used as loading control. ** and * indicate potentially nonspecific bands. (C) Immunohistochemistry shows Kdm3a localization in acrosomes (ac) of wild-type (WT) mature spermatids but not Kdm3a∆JC/∆JC as detected with C-Kdm3a antibody described in (B). Arrows indicate the dorsal surface of the spermatid on DAPI-stained nuclei.

Reduced methylation state of Hsp90 is observed in Kdm3a∆JC/∆JC pull downs relative to wild-type (WT) extracts. This difference is not detectable in shorter exposures of hsp90aa1 blots (unpublished data). (D) Fold change in the methylation levels of histones as detected with a monoclonal antibody to dimethylated H3K9. Each point represents the ratio between the mean intensity of H3K9 signal in transfected cells relative to their neighboring untransfected cells. The error bars represent the SEM between three independent experiments. More than 100 cells were counted for each condition. (E) Endogenous levels of Kdm3a protein are increasingly reduced as Hsp90 is inhibited with 17AAG in RPE1 cells. Tubulin is used as a loading control. (F) Box plots illustrate the subcellular distribution of RFP-Kdm3a isoforms in cells grown at the indicated 17AAG dose. Maximal pixel intensity of the nuclear and cytoplasmic compartments was measured as indicated in Figure S1C. * indicates statistical significance (p < 0.001, t test). (G) Change of subcellular distribution of Kdm3a-i2 after Hsp90 inhibition. Note nuclear depletion (red arrow) is often accompanied by increased H3K9 methylation (green arrow). Scale bars: 20 μM.
components of microtubule and actin networks in animals lacking Kdm3a demethylase activity. The differential state of cellular chaperones in Kdm3aΔ/ΔC/Lc samples could reflect the state of actin or other folding targets altered in Kdm3a mutants.

**DISCUSSION**

In this study, we begin to unravel the functional complexity of Kdm3a. We present genetic and biochemical evidence that cytoskeletal abnormalities contribute to the male infertility phenotype previously reported for Kdm3a mouse models. The use of two Kdm3a models has allowed us to separate cytoplasmic from chromatin condensation defects. The most severe knockout model has acroplaxome, manchette, and HTCA abnormalities accompanying variable states of DNA condensation. In the milder gene-trap model, DNA condenses fairly normally, but manchette and centriolar defects persist, and mature spermatids still have rounded heads. The variable penetrance of these traits perhaps reflects the different dependence of multiple processes on Kdm3a activity. The Kdm3a isoform newly identified here may contribute to phenotype amelioration in the second mouse model. We further demonstrate that Kdm3a interacts with and requires Hsp90 for its demethylation activity and protein stability. We show that Kdm3a localizes not only to the nucleus but also to different cytoplasmic compartments depending on growth conditions and Hsp90 activity. We found Kdm3a to be enriched in the acrosome, which is in turn disrupted in Kdm3aΔ/ΔC/Lc mutants and provide evidence that the state of actin and tubulin components that form these structures is altered, although their protein levels remain unchanged. We therefore propose that Kdm3a is directly involved in the cytoskeletal rearrangements required to produce mature spermatids. Whether this is through Kdm3a interactions with actin, tubulin, or dynein components identified as candidates in this study or their postranslational modification (PTM) remains to be investigated.

The original finding of the dynamic distribution of Kdm3a in the cytoplasm of spermatids (Okada et al., 2007) suggested that Kdm3a could have a cytoplasmic function. Hsp90 chaperones have also been reported among Kdm3a pulled-down proteins from somatic cells (Brauchle et al., 2013). Our ultrastructural analysis revealed cytoskeletal defects known to depend on actin, tubulin, and cellular chaperones (Kierszenbaum et al., 2011); biochemical evidence then uncovered their abnormal partition after Kdm3a loss of function. This could be a consequence of altered functional capacity of cellular chaperones, known to regulate protein folding and to be dynamically methylated (Abu-Farha et al., 2011), but the methylation state of Hsp90 was unchanged in Kdm3a homozygous mice. Instead, we found that Kdm3a protein stability and enzymatic activity is highly dependent on Hsp90, raising the possibility that the observed cytoskeletal defects could therefore be a direct consequence of Kdm3a activity on cytoplasmic components for which it would require Hsp90 chaperoning.

Actb2 is differentially enriched in Kdm3aΔ/ΔC/Lc samples purified with antibodies to methyl lysines and Hsp90. Elongation of the spermatid head requires extensive restructuring of actin filaments in the leading edge of the acrosome and nuclear envelope (Kierszenbaum et al., 2003). Changes in the actin network are driven by a large variety of biochemical properties that involve nucleation of monomers and elongation of filaments to produce the many distinct actin architectures observed in living organisms (Blanchoin et al., 2014). Biochemical evidence suggests that the properties of actin filaments may vary according to the mix of actin isoforms in the filament (Perrin and Ervasti, 2010). If Actb2 is postranslationally altered in Kdm3a mutant spermatids, its association with other actin monomers may perturb filament formation and disrupt acrosome development, giving rise to the abnormal acrosome observed in Kdm3aΔ/ΔC/Lc mice. The importance of the correct spatiotemporal sequence in actin dynamics is illustrated by the many disease states associated with altered actin cytoskeleton (Blanchoin et al., 2014). Indeed, Actb2 is associated with hepatocellular carcinomas (Chang et al., 2011; Junrong et al., 2011). Interestingly, Kdm3a may play a significant role during the malignant transformation of hepatocellular carcinomas (Yamada et al., 2012). Furthermore, this study found that in 80% of the samples expressing Kdm3a protein, this protein localized exclusively to the cytoplasm (Yamada et al., 2012). Actb2 is therefore a very strong candidate to molecularly underlie Kdm3a mutant acrosomes, and this functional interaction may extend to somatic cells. Cytoplasmic dyneins are also associated with microtubules of the manchette for transport (Yoshida et al., 1994). These proteins found to be lysine methylated in human cells (Hornbeck et al., 2012; Cao et al., 2013) are thus also potential candidates via which Kdm3a could impact the cytoplasmic aspect of spermatogenesis found altered in Kdm3a mutant mice.

None of the proteins analyzed in this study shows altered levels in Kdm3aΔ/ΔC/Lc samples. Likewise, publicly available microarrays in which Kdm3a has been silenced (Hertzog et al., 2012) show no significant change in the transcript levels of Actb2, dyneins, or Ccts, indicating the impact of Kdm3a on these cytoplasmic components is not at the transcriptional level. The effect on these components could thus be through direct protein interaction and/or postranslational modifications. The functional requirement of Hsp90 for Kdm3a demethylase activity raises the possibility that Kdm3a cytoplasmic targets could be postranslationally modified as they are being folded, with the modifications contributing to their tertiary structure or polymerization state. The consistent lack of the ventral surface of the acrosome in Kdm3a mutant mice is intriguing. Could postranslational...
modifications act to demarcate spatial boundaries between dorsal and ventral acrosome surfaces in otherwise continuous filaments of actin. Posttranslational modifications of ubiquitous and abundant proteins may thus confer spatial compartmentalization. Another possibility is that the ventral and dorsal surfaces of the acrosome could contain different ratios of actin isoforms.

Only a small fraction of cellular proteins are folded by Cct chaperonin, and from these only 5–10% will require further interaction with Hsp90 chaperones (Kim et al., 2013). It is thus likely that Kdm3a interacts with the Hsp90 chaperone system independently from chaperonins fulfilling other functions not directly related to protein folding. Histone and nonhistone substrates for Kdm3a are not mutually exclusive scenarios and warrant further investigation. We analyzed only one methylated residue of Hsp90, but it is known to have at least two methylatable residues (Abu-Farha et al., 2011); the effect of Kdm3a on these alternative sites remain to be investigated. The methylation state of Cct chaperonins also deserves further investigation.

Hsp90 and Cct chaperones are also known to play a role in chromatin remodelling by modulating binding and release of transcriptional regulatory machineries (Liu and DeFranco, 1999; Freeman and Yamamoto, 2001) and binding to heterochromatin in germ cells (Souèes et al., 2003). We investigated the effect of Hsp90 on Kdm3a histone demethylase activity and found that it largely depends on these chaperones. Interestingly, the enzymatic activity, protein stability, and subcellular localization of Kdm3a depended on Hsp90. Further studies would be particularly relevant for androgen-responsive genes known to be regulated both by Hsp90 and Kdm3a (Freeman and Yamamoto, 2001; Yamane et al., 2006) and for the cellular response to hypoxia in which Kdm3a and HIF1-α levels are regulated by oxygen (Beyer et al., 2008; Pollard et al., 2008; Wellmann et al., 2008) and Hsp90 (Minet et al., 1999), respectively. Interestingly, we also found the JmJC domain of Kdm6a to interact with Hsp90aa1 (Figure S6), suggesting that interaction between JmJC-containing proteins and Hsp90 chaperone machinery is a common feature and may be relevant in the enzymatic activity of this family in general.

In summary, our work has only just begun to illustrate the complex regulatory and spatial landscape of Kdm3a, providing evidence for a wider functional role of this lysine demethylase beyond histone modification in the nucleus.

**MATERIALS AND METHODS**

**Kdm3a mouse models**

*Kdm3a*Δ*JC* mice (*Kdm3a*Δ*JC* ES cells (a gift from Y. Zhang, Harvard Medical School) were used to produce chimeric animals upon microinjection into C57BL6J blastocysts and to establish a parental Kdm3a colony. Heterozygous males were then crossed to C57BL6 females expressing ubiquitous Cre, and the resulting mice containing recombination Kdm3a alleles (*AJC*) were identified by PCR. LacZ positive, F1 mice were crossed with a ubiquitously expressing Cre line to produce knockout offspring lacking the JmJC catalytic domain. Subsequent generations were backcrossed to the C57BL6 strain and genotyped with primers A, B, and C (Tateishi et al., 2009) listed in Table S1.

*Kdm3a*GT mice were generated by microinjection of XR0062 ES cell lines (129/ola), obtained from the Sanger Institute Gene Trap Resource (www.genetrap.org), into C57BL6J blastocysts with transmission of the mutation on a C57 background. The cells used contained a gene-trap insertion (*Kdm3a*GT) within intron 5–6 of the mouse Kdm3a locus. Genotypes were identified by PCR of genomic DNA with primers GT1, GT2, and GT3 described in Table S1. Expression levels of the full-length transcript in gene-trapped mice were determined with the primer pair GT4.

**Antibodies**

The antibodies used along this study are to KDM3A (12835; ProteinTech; and NB100-77282; Novus Biologicals, Littleton, CO); KDM3B (HIC 00189; Bethyl Laboratories, Montgomery, TX); Cct4 (ARP34271; Aviva); anti–mono- and dimethylated lysines (ab23366 and ab76118; Abcam, Cambridge, MA); anti-GFP (sc-8334), monoclonal to GAPDH (S019A-2; Imagegex, San Diego, CA), β-gal (A-11132; Molecular Probes), GST (C83271; LSBio); HP1α (clone 15.1952; Upstate); γ-tubulin (GTU-88; Sigma-Aldrich, St. Louis, MO), β-actin (ab8229; Abcam), Hsp90ab1 (MAB32861; R&D Systems), Hsp90aa1 (10713715; Pierce, Rockford, IL), Actbl2 (ab134977; Abcam). Secondary antibodies: anti-mouse and anti-rabbit F(ab′)2 immunoglobulin G Alexa Fluor 488 and 584 (Molecular Probes) for immunofluorescence or horseradish-conjugated peroxidase for immunoblots.

**Cell lines, transfections, and cell fractionation**

Telomerase-immortalized human retinal pigment epithelial cells (hTERT-RPE1) were obtained from Clontech (Mountain View, CA). Transfections were done in RPE1 cells at 80% confluency and were seeded the day before transfection without antibiotic. Lipofectamine 2000 was used following the manufacturer’s guidelines for 5 h. MEFs were isolated by mincing E11.5- to E13.5-d embryos in DMEM supplemented with antibiotics and serum. MEFs were grown in 3% O2 incubators. Subcellular fractionation of MEFS was performed using Qproteome Cell Compartment kit according to the manufacturer’s instructions (Qiagen, Valencia, CA).

**Primers, RT-PCR, and qRT-PCR**

All primers used below are listed in Table S1. First-strand cDNA was synthesized with random primers and Superscript III (Invitrogen, Carlsbad, CA). qPCRs were set up using the Quantitect SYBR green PCR kits (Qiagen). Individual reactions totaled 10 μl with a final primer concentration of 0.5 mM.

**Cloning of Kdm3a-i2 isoform**

*Kdm3a*–*i2* was amplified from homozygous Kdm3a-GT MEF RNA or wild-type testis RNA and reverse transcribed with random primers; this was followed by PCR with primers designed from ENSEMBL prediction ENSMUST00000101304 as listed (Table S1). The PCR product was cloned into pGem-Teasy (Promega, Hayward, CA) vector and sequenced to verify correct splicing and open reading frame.

**Tagged constructs**

Tagged constructs were made using cDNA from homozygous Kdm3a-GT MEF RNA or wild-type testis RNA and reverse transcribed with random primers; this was followed by PCR with primers designed from ENSEMBL prediction ENSMUST00000101304 as listed (Table S1). The PCR product was cloned into pEGFP or RFP-vector. Kdm3a-FL was subcloned in-frame into pEGFP or RFP vector. Kdm3a-FL was subcloned in-frame into RFP vector. Primers contained Xhol and BamHI restriction sites (Table S1).

**TEM**

Adult mice testes were fixed in 2% paraformaldehyde, 2.5% glutaraldehyde in 0.1 M sodium cacodylate buffer plus 0.04% CaCl₂ overnight. Several cuts at the edges of the testes were done to allow penetration of fixative.

**Immunofluorescence**

Mouse testes utilized for tissue sections were fixed in ice-cold 4% methanol-free formaldehyde/phosphate-buffered saline (PBS) for at least 1 h (formaldehyde from TAAB Bioscience). Fixed material was dehydrated in methanol, wax embedded, and microtome sectioned.
In all cases, cells and sections were incubated with primary antibodies overnight at 4°C. Secondary antibody was incubated for 2 h at room temperature or overnight at 4°C. Investigation of Kdm3a presence in the cytoplasm of spermatids was done on dissected tubules (Kotaja et al., 2004) squashed on glass slides and frozen on dry ice; this was followed by methanol and acetone fixation (Zhou et al., 2009).

Protein extracts and immunoprecipitations

Total cell extracts were obtained by lysing cells in 150 mM Tris-HCl (pH 7.5), 150 mM NaCl, 0.5% octylphenoxypolyethoxyethanol (IGEPAL), 1% Triton, 1 mM EDTA, 0.5% deoxycholate with Complete EDTA-free protease inhibitors (Roche, Indianapolis, IN), and Benzoaze DNaSe (Novagen). For immunoprecipitations, cells were lysed in 50 mM Tris-HCl (pH 7.5), 100 mM NaCl, 10% glycerol, 5 mM EDTA, and 0.5% IGEPAEl with Benzoaze DNaSe and protease inhibitors. Successive washes were carried out with reducing IGEPAL content to a final concentration of 0.01%, with final washes using only Tris-HCl, NaCl, EDTA, and 10% glycerol. Primary antibody was generally incubated overnight at 4°C, this was followed by 45 min with Dynabeads (Invitrogen) to concentrate immunoglobulin complexes. Hsp90 pull downs were done as previously described (Taipale et al., 2012). For the identification of methylated peptides, proteins extracts were prepared in the presence of 160 μM of the demethylase inhibitor 5-carboxy-8-hydroxyquinoline (SML0067; Sigma-Aldrich).

Sucrose gradients

Testis cytoplasmic extracts were prepared by hypotonic shock. Briefly, gently dislodged testes were taken to single-cell suspension in ice-cold PBS by passaging them 10 times through a 1-ml plunger. After being rinsed twice in PBS, cells underwent 5 min of centrifugation at 1000 × g. Cells were suspended in 0.1 × TBS (50 mM Tris, pH 7.5, 150 mM NaCl)–8% sucrose and left until they became swollen (for 30–60 min) on ice. Swollen cells were centrifuged at 10 min of centrifugation at 2500 × g. All supernatant was removed, and cells were resuspended in 152 μl of 8% sucrose and 1.37 ml of lysis buffer (1 mM HEPES, pH 7.2, 0.5% IGEPAEl, 0.5 mM MgCl2, 0.1% β-mercaptoethanol, Complete EDTA-free protease inhibitor); this was followed by 10 min of centrifugation at 2500 × g. An aliquot of this cleared cytoplasmic extract represented gradient input; the remaining extract (1 ml approximately) was layered on top of a discontinuous sucrose gradient formed by 1 ml of 40%, 1 ml of 50%, and 2 ml of 70% sucrose layers in gradient buffer (10 mM PIPES, pH 7.2, 0.1% Triton X-100, 0.1% β-mercaptoethanol). Gradients were run overnight at 40,000 rpm in a Beckman ultracentrifuge. Fractions of 250 μl were collected manually from the top of the gradients. Nocodazole (10 μg/ml) and cytochalasin B (5 μg/ml) were added to testis extracts in ice-cold PBS by passaging them 10 times through a 1-ml plunger.

MS

For analysis of gel fractions, pulled-down proteins were fractionated on an SDS–PAGE gel and stained with TrueBlue (E gedon). The indicated gel chunks were excised and cut into 1- to 10-mm cubes; this was followed by in-gel digestion using a ProGest Investigator in-gel digestion robot (Genomic Solutions, Ann Arbor, MI) using standard protocols (Shevchenko et al., 1996). These gel cubes were destained by washing with acetonitrile and subjected to reduction and alkylation before digestion with trypsin at 37°C. Peptides were extracted with 10% formic acid and concentrated down to 20 μl using a SpeedVac (ThermoSavant, Thermo Fisher Scientific, Lafayette, CO). The peptides were then separated using a NanoLC Ultra 2D Plus loading pump and NanoLC AS-2 autosampler equipped with a nanoflex chiPLC chip-based chromatography system (Eskigent, Redwood City, CA) and a ChromXP C18-CL trap and column (Eskigent), and eluted with a gradient of increasing acetonitrile containing 0.1% formic acid (5–35% acetonitrile in 45 min, 35–50% in a further 3 min, followed by 95% acetonitrile to clean the column, before reequilibration to 5% acetonitrile). The eluent was sprayed into a TripleTOF 5600 electrospray tandem mass spectrometer (ABSciex, Foster City, CA) and analyzed in information-dependent acquisition mode; we performed 250 ms of MS followed by 100-ms tandem mass spectrometry (MS/MS) analyses on the 20 most intense peaks seen by MS. The MS/MS data file generated was analyzed using the ProteinPilot (version 4.5 beta) Paragon algorithm (ABSciex) against the SwissProt database (www.ebi.ac.uk/uniprot, accessed February 2013) with both no species restriction and mouse-only, trypsin as the cleavage enzyme, and carbamidomethyl modification of cysteines. For bead analysis, pulled-down proteins were digested on beads at 37°C overnight using trypsin without prior reduction or alkylation. The peptides were then desalted using C18 Zip Tips (Millipore, Billerica, MA) and loaded onto a PepMap100 RSLC nanotrap column (Thermo Fisher Scientific) at a flow rate of 1 μl/min. Peptides were separated on a 15-cm PepMap100 RSLC analytical column (Thermo Fisher Scientific) using a Dionex Ultimate RSLC nano system and eluted with a gradient of increasing acetonitrile containing 0.1% formic acid (1.6–32% acetonitrile in 45 min, 32–90% in a further 0.1 min, followed by 4.6 min at 90% acetonitrile and back to 1.6% to re-equilibrate the column). Peptides were eluted into a Q-Exactive mass spectrometer (Thermo Fisher Scientific) using a flow rate of 300 nl/min and a spray voltage of 4 kV. Proteome Discoverer (version 1.4) was used to create peak lists from raw data and to search against the SwissProt Mus musculus database (accessed November 2013). Search parameters were set as follows: precursor mass tolerance of 10 ppm, fragment ion mass tolerance to 0.03 mDa, and enzyme as trypsin, allowing two missed cleavages. No static modifications were specified. However, the dynamic modification of single, di-, and tri-methylation was allowed. Data were then further analyzed using Scaffold 4 (version 4.2.1).

Two-hybrid screens and GST subcloning

A mouse cDNA library was prepared by reverse transcribing mouse embryonic eye RNA with oligo-dT primers containing a BamHI site and ligated to EcoRI adaptors. cDNA products were digested with BamHI/EcoRI and cloned onto BamHI/EcoRI sites of pGBK7T (Invitrogen) previously linearized with CcrI. The generated library was transformed by electroporation to a pJ69-a yeast strain. The indicated region of Hsp90ab1 was cloned onto a pGADT7 bait vector, and this transformed into the mating-compatible strain AH109-a. Strains were mated as described by the vector manufacturer (Invitrogen). Interacting proteins were identified by growing mated cultures in triple-selection media lacking adenine, leucine, and tryptophan. Larger colonies were assessed for X-gal staining, and tryptophan. Larger colonies were assessed for X-gal staining, and positive clones were streaked onto new triple-selection media. cDNA from grown colonies was then amplified by PCR and sequenced with pGADT7 primers (Invitrogen). DNA was purified from selected colonies and retransformed onto yeast for further validation. For GST constructs, isolated plasmids were transformed into bacteria, amplified, and digested with EcoRI/XhoI to clone onto pGEX-4-T1. Hsp90a1ΔC and Hsp90a1-C terminus baits are in pAS2-1 vectors (Obermann et al., 1998).
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