Phosphorylation of the RNA-binding protein Dazl by MAPKAP kinase 2 regulates spermatogenesis

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ABSTRACT Developing male germ cells are exquisitely sensitive to environmental insults such as heat and oxidative stress. An additional characteristic of these cells is their unique dependence on RNA-binding proteins for regulating posttranscriptional gene expression and translational control. Here we provide a mechanistic link unifying these two features. We show that the germ cell–specific RNA-binding protein deleted in azoospermia-like (Dazl) is phosphorylated by MAPKAP kinase 2 (MK2), a stress-induced protein kinase activated downstream of p38 MAPK. We demonstrate that phosphorylation of Dazl by MK2 on an evolutionarily conserved serine residue inhibits its interaction with poly(A)-binding protein, resulting in reduced translation of Dazl-regulated target RNAs. We further show that transgenic expression of wild-type human Dazl but not a phosphomimetic form in the Drosophila male germ line can restore fertility to flies deficient in boule, the Drosophila orthologue of human Dazl. These results illuminate a novel role for MK2 in spermatogenesis, expand the repertoire of RNA-binding proteins phosphorylated by this kinase, and suggest that signaling by the p38-MK2 pathway is a negative regulator of spermatogenesis via phosphorylation of Dazl.

INTRODUCTION
Spermatogenesis is a highly coordinated process during which developing male germ cells experience periods of transcriptional quiescence despite a requirement for continued protein synthesis (Hecht, 1998). The process is therefore heavily reliant on posttranscriptional mechanisms for regulating gene expression. RNA-binding proteins fulfill this function by stabilizing presynthesized transcripts that are to be translated at later stages of spermatogenesis (Paronetto and Sette, 2010; Idler and Yan, 2012). Consequently, deletion of several RNA-binding proteins in male mice results in meiotic arrest and infertility (Venables and Cooke, 2000; Dass et al., 2007; Gutti et al., 2008; Hsu et al., 2008), illustrating the importance of these proteins in male germ cell development. An additional property of developing male germ cells is their exquisite sensitivity to heat stress, and even mild, transient hyperthermia induces germ cell apoptosis with detrimental consequences for fertility (Yin et al., 1997; Rockett et al., 2001). Although the underlying molecular mechanisms that mediate these effects are not entirely understood, p38 MAP kinase plays an important role in this process (Jia et al., 2009). A primary target of activated p38 is the serine/threonine kinase mitogen-activated protein kinase–activated protein kinase 2 (MK2/MAPKAPK2; Stokoe et al., 1992). Direct phosphorylation of MK2 on two threonine residues (T222 and T334) results in its activation (Ben-Levy et al., 1995). However, the precise role of MK2 in spermatogenesis is unknown.

Here we show that the germ cell–specific RNA-binding protein deleted in azoospermia-like (Dazl) is phosphorylated by MK2. We demonstrate that phosphorylation of Dazl by MK2 inhibits its association with poly(A)-binding protein, resulting in reduced translation of Dazl-regulated RNA targets both in vitro and in a transgenic Drosophila model system. These findings establish a novel connection between stress-induced kinase signaling and translational regulation by RNA-binding proteins in the male germline.

RESULTS AND DISCUSSION
Dazl is a novel substrate of MK2
To characterize a potential role for MK2 signaling in male germ cell development, we conducted a proteomic screen to identify novel substrates of this kinase. Expression of MK2 was reduced by short hairpin RNA (shRNA) in SUM149 cells, a triple-negative breast

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Abbreviations used: Dazl, deleted in azoospermia-like; MK2/MAPKAPK2, mitogen-activated protein kinase–activated protein kinase 2; PABP, poly(A)-binding protein; RRM, RNA recognition motif.
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Dazl is a novel substrate of MK2. (A) Reduction of MK2 expression by shRNA. Cell lysates from SUM149 cells stably expressing either control or MK2 shRNA were resolved by SDS–PAGE (on two separate gels) and immunoblotted with the indicated antibodies. (B) Schematic of the screen used to identify novel MK2 substrates. Lysates from control and MK2 shRNA cells were incubated with human protein microarrays and probed with anti-phosphoserine and anti-phosphothreonine antibodies, followed by detection with Alexa 488–conjugated secondary antibodies. Arrays were scanned and quantified. See Materials and Methods for additional details. (C) Bar graph of the top 20 putative MK2 substrates identified from the screen. Data are plotted as the average fold signal reduction obtained from the MK2-knockdown shRNA array compared with the control shRNA knockdown array. Error bars indicate SD. (D) In vitro kinase assay using GST-Dazl and MK2. GST-Dazl (or no substrate) was incubated in the presence or absence of active recombinant MK2 and ATPγS, followed by treatment with PNBM. Kinase reactions were resolved by SDS–PAGE (on three separate gels), followed by immunoblotting with the indicated antibodies.

Among the top 20 putative MK2 substrates (Figure 1C) was deleted in azoospermia-like (Dazl), a germ cell-specific RNA-binding protein. Dazl belongs to the DAZ family of proteins, which also includes DAZ and Boule (Brook et al., 2009). The DAZ family of proteins has an evolutionarily conserved role in regulating gametogenesis (Xu et al., 2001; Reynolds and Cooke, 2005; Vangompel and Xu, 2011). DAZL-knockout male mice are sterile due to a block in meiosis (Ruggiu et al., 1997; Saunders et al., 2003), and men with mutations in this RNA-binding protein exhibit varying degrees of spermatogenic defects (Tung et al., 2006). The fact that Dazl was identified in this screen was of particular interest, given the established function of MK2 in phosphorylating and regulating several other RNA-binding proteins in the context of inflammation (Gaestel, 2006). MK2, for instance, phosphorylates the RNA-binding proteins...
MK2 phosphorylates Dazl at S65 within the RNA recognition motif

To investigate the biological role of this posttranslational modification, we determined the residue(s) on Dazl that are phosphorylated by MK2. Dazl consists of two functional domains (Figure 2A): an evolutionarily conserved RNA recognition motif (RRM), which binds target RNA sequences, and a deleted in azoospermia (DAZ) repeat important for protein–protein interactions (Brook et al., 2009). The consensus motif for MK2 phosphorylation has been determined (Stokoe et al., 1993), and its defining features include the presence of an arginine residue at the −3 position (i.e., three residues N-terminal to the phosphoacceptor site) and hydrophobic residues at the −5 and −6 positions. Primary sequence analysis of Dazl revealed a serine residue (S65) in the RRM located within an ideal consensus motif for MK2 phosphorylation (Figure 2A). To test whether this site was phosphorylated by MK2, we generated GST-fusion proteins of either the wild-type RRM or a mutant RRM in which this serine residue was replaced with nonphosphorylatable alanine (S65A). These fusion proteins were used as substrates in in vitro kinase reactions with active recombinant MK2 (Figure 2B). The results demonstrate that S65 is a site of MK2 phosphorylation, as mutation of this residue to alanine abolished the signal. To test whether S65 is the sole site of MK2 phosphorylation, we produced full-length GST-Dazl and GST-Dazl-S65A and used them as substrates in in vitro kinase assays. Results of this experiment reveal that mutation of this residue within the context of the full-length protein abrogated the signal, indicating that S65 is the only residue in Dazl phosphorylated by MK2 (Supplemental Figure S1). Of interest, S65 and its surrounding amino acid sequence are conserved in Dazl from zebrafish and mouse as well as in human DAZL (Xu et al., 2001), and S65 phosphorylation of murine Dazl has been detected by mass spectrometry (Wu et al., 2012), indicating that this posttranslational modification occurs in vivo.

To study the physiological relevance of Dazl-S65 phosphorylation, we generated a phospespecific antibody. We first validated the specificity of this reagent by performing an in vitro kinase assay using MK2 and GST-Dazl-RRM or GST-Dazl-RRM-S65A. A signal was detected when MK2 was incubated with the wild-type Dazl-RRM but not Dazl-RRM-S65A (Figure 2C), demonstrating that the antibody detects Dazl only when phosphorylated at S65. To determine whether phosphorylation of Dazl-S65 could be detected in cells, we cotransfected HEK293 cells with constructs encoding hemagglutinin (HA)-Dazl and myc-MK2. Cells were then stimulated (or not) with sodium arsenite, a chemical that induces oxidative stress and is a potent activator of p38 MAPK. Immunoblotting of cell lysates demonstrated activation of p38 and MK2 upon stimulation and a signal using the anti–phospho-Dazl-S65 antibody (Figure 2D).

We then addressed whether genetic perturbation of the p38-MK2 signaling pathway affects phosphorylation of Dazl. HEK293 cells were transfected with plasmids encoding HA-Dazl, myc-MK2, and either a dominant-negative mutant of p38α (FLAG-p38-MAF) or a constitutively active form of MK2 (myc-MK2-EE). Immunoblotting of cell lysates revealed a lack of Dazl-S65 phosphorylation in arsenite-stimulated cells expressing dominant-negative p38 (Figure 2D). However, even in the absence of stimulation, Dazl-S65 phosphorylation was detected in cells expressing constitutively active MK2 (Figure 2D). Furthermore, chemical inhibition of the p38-MK2 pathway using the p38 MAPK inhibitor SB203580 or MK2 inhibitor III confirmed that phosphorylation of Dazl-S65 was dependent on MK2 kinase activity (Figure 2E). Collectively these data indicate that serine 65 of Dazl is phosphorylated by MK2.

S65 phosphorylation of Dazl negatively regulates translation but not RNA binding

Within male germ cells, Dazl stabilizes several mRNAs that encode proteins with critical functions in meiosis, such as the synaptonemal complex component Syce3 (Reynolds et al., 2007) and the DEAD-box RNA helicase Mvh/Ddx4 (Reynolds et al., 2005). As a result, deletion of DAZL in male mice results in reduced protein expression of Syce3 (Reynolds et al., 2007) and Mvh (Reynolds et al., 2005), culminating in meiotic arrest.

Previous studies examining the RNA-binding properties of Dazl determined that the protein preferentially binds to a loosely defined uridine-rich consensus sequence in the 3′ untranslated region (UTR) of these transcripts (Venables et al., 2001; Jiao et al., 2002; Magee et al., 2002). Because the MK2 phosphorylation site we identified was located within the RRM of Dazl, we speculated that modification at this site affects RNA binding. To directly test this hypothesis, we used surface plasmon resonance (SPR). Synthetic biotinylated RNA molecules consisting of the Dazl-binding consensus motif (5′-UUCGU3′) or a nonspecific mutant RNA molecule (5′-A6C6A6CA6CA6A-3′) were immobilized on a NeutrAvidin-coated sensor chip. Dazl-RRM-WT or S65D proteins were flowed over the chip as analytes, and binding affinities were determined. The results demonstrate that although neither protein binds to the nonspecific RNA, both the wild-type RRM and the S65D mutant RRM bind to the RNA consisting of the Dazl-binding consensus sequence (Figure 3A). Furthermore, determination of the dissociation constant (Kd) similarly revealed no significant difference in the association of either protein with the Dazl-binding RNA consensus sequence (Figure 3B and Supplemental Figure S2). These findings are consistent with the crystal structure of the murine Dazl RRM, which was solved in complex with three different target RNAs (Jenkins et al., 2011) and in which it is demonstrated that S65 does not make direct contact with the RNA molecule. These results therefore suggest that posttranslational modification of this residue may regulate an alternative function of Dazl.

In response to heat stress in germ cells, Dazl localizes to stress granules (Kim et al., 2012)—nonmembranous cytoplasmic foci composed of stalled translational complexes and associated RNA-binding proteins. In the absence of Dazl, stress granules do not form, and germ cells are more susceptible to stress-induced apoptosis (Kim et al., 2012). We therefore tested whether mutation of S65 affects localization of Dazl to stress granules in response to arsenite treatment (Supplemental Figure S3). The results demonstrate that both wild-type and mutant forms of Dazl localize to stress granules, suggesting that modification of this residue is likewise not involved in mediating localization to these structures.

Once bound to the 3′ UTR of target mRNAs, Dazl functions as a translational enhancer by recruiting poly(A)-binding protein (PABP) to stimulate the initiation of translation (Collier et al., 2005; Brook et al., 2009). We therefore postulated that phosphorylation of S65 may affect translation by altering the association of Dazl with PABP.
FIGURE 2: MK2 phosphorylates Dazl at S65. (A) Domain architecture of human Dazl. DAZ, deleted in azoospermia domain; RRM, RNA recognition motif. The putative MK2 phosphorylation site (S65) is indicated in red. (B) MK2-mediated phosphorylation of Dazl-S65 in vitro. Wild-type or S65A GST-Dazl-RRM (residues 32–117) fusion proteins were incubated with active recombinant MK2 and ATPγS, followed by incubation with PNBM. Kinase reactions were resolved by SDS–PAGE (on two separate gels), followed by immunoblotting with the indicated antibodies. (C) Detection of Dazl-S65 phosphorylation using a phosphospecific antibody. GST-Dazl RRM (wild-type or S65A) fusion proteins were incubated with active recombinant MK2 and ATP. Kinase reactions were resolved by SDS–PAGE (on two separate gels), followed by immunoblotting with the indicated antibodies. (D) Alteration of Dazl-S65 phosphorylation status via genetic manipulation of the p38-MK2 signaling pathway. HEK293 cells were transfected with the indicated constructs and either stimulated or not with sodium arsenite (Ars) and lysed. Cell lysates were split, resolved by SDS–PAGE (on eight separate gels), and immunoblotted with the indicated antibodies. Western blots shown are representative of three independent experiments. (E) Reduced Dazl-S65 phosphorylation by chemical inhibition of the p38-MK2 signaling pathway. HEK293 cells were transfected with the indicated constructs and pretreated for 30 min with DMSO (vehicle) only, the p38 inhibitor SB20350 (10 μM), or MK2 inhibitor III (20 μM). Cells were then stimulated with sodium arsenite and lysed. Cell lysates were split, resolved by SDS–PAGE (on seven separate gels), and immunoblotted with the indicated antibodies. Western blots shown are representative of three independent experiments.
and we addressed this possibility using a luciferase reporter assay. A plasmid was constructed in which the 3′ UTR of murine SYCP3, a validated Dazl-binding sequence (Reynolds et al., 2007), was cloned downstream of the Renilla luciferase gene. HeLa cells were then transfected with this construct or cotransfected with this construct and a plasmid encoding myc-Dazl-WT, myc-Dazl-S65D, or, as a negative control, myc-Dazl-R115G, a mutant that is defective in RNA binding and is associated with human infertility (Tung et al., 2006; Jenkins et al., 2011). Western blotting was used to confirm equal expression of the myc-tagged proteins (Figure 3C), and luciferase
activity was measured to assay translation. The results demonstrate significantly less luciferase activity with expression of Dazl-R115G than with wild-type Dazl (Figure 3D). Moreover, cells expressing Dazl-S65D also displayed decreased luciferase activity (Figure 3D), suggesting that while mutation of this residue does not affect RNA binding, it does decrease translation. To determine whether this reduction in translation is due to inhibition of PABP binding, we co-transfected HEK293 cells with plasmids encoding untagged PABP and myc-Dazl-WT, myc-Dazl-S65D, myc-Dazl-S65A, or empty vector as a control. PABP was immunoprecipitated from cell lysates, and the immunoprecipitates were probed for Dazl (Figure 3E). Less Dazl-S65D coprecipitated with PABP than with Dazl-WT (and, reciprocally, less PABP coprecipitated with Dazl-S65D than with Dazl-WT; Supplemental Figure S4), whereas expression of Dazl-S65A appeared to stabilize the interaction (Figure 3E and Supplemental Figure S4). Taken together, these results suggest that MK2-mediated phosphorylation of Dazl decreases translation by inhibiting the interaction of Dazl with PABP.

Expression of human Dazl, but not Dazl-S65D, restores fertility in boule mutant flies

The Drosophila orthologue of Dazl is Boule, and boule mutant male flies are sterile due to a defect in meiotic entry (Eberhart et al., 1996). Specifically, spermatocytes of boule mutants are arrested in meiosis I at the G2/M transition due to reduced translation of Twine (Maines and Wasserman, 1999), the meiotic Cdc25 phosphatase that dephosphorylates the cdc2/cyclinB complex. Overexpression of Twine in the boule mutant genetic background rescues this phenotype (Maines and Wasserman, 1999), as does transgenic expression of either human Boule (Xu et al., 2003) or Xenopus Dazl (Houston et al., 1998), illustrating the high degree of functional conservation among DAZ family members.

On the basis of these studies, we predicted that expression of human Dazl would rescue the Drosophila boule phenotype, whereas Dazl-S65D would not due to its reduced capacity to promote translation. To directly test this, we expressed myc-tagged human Dazl or Dazl-S65D in the male germline of Drosophila mutant genetic background (Figure 4A). The fertility of male flies expressing these transgenes was assessed by a fertility assay. Expression of wild-type Dazl restored fertility in the boule mutant genetic background (Figure 4B), indicating that human Dazl can functionally compensate for the lack of Drosophila boule. Furthermore, analysis of testes by phase contrast microscopy revealed rescue of meiotic entry based on the presence of postmeiotic round spermatids and elongating sperm bundles in flies expressing wild-type Dazl (Figure 4C). In contrast, germ cells from flies expressing Dazl-S65D did not differentiate beyond the spermatocyte stage and phenotypically resembled boule mutants, demonstrating that the S65D mutation impairs the function of Dazl in vivo.

To determine whether the rescue of meiotic entry and restoration of fertility upon expression of wild-type Dazl correlated with restored expression of Twine, we analyzed levels of Twine protein in testes from flies of the aforementioned genotypes. Consistent with previous studies, we observed reduced levels of Twine in boule mutant testes that were restored upon expression of wild-type Dazl (Figure 4D). In contrast, expression of Dazl-S65D did not rescue Twine expression in the boule mutant genetic background (Figure 4D). Taken together, these results provide evidence that the phosphorylation status of Dazl-S65 regulates Twine translation, meiotic progression, and male fertility.

In summary, we found that phosphorylation of the RNA-binding protein Dazl by MK2 is a negative regulator of male meiosis (Figure 4E). Our data suggest that this is due to decreased recruitment of PABP, ultimately resulting in reduced translation of Dazl-regulated target RNAs. The precise molecular mechanism by which this occurs is unclear, although we speculate that it might involve a conformational change, since S65 of Dazl lies outside of the minimal region required for the PABP-Dazl interaction (Collier et al., 2005).

An intriguing aspect of this work is the regulation of meiotic G2/M progression by MK2-mediated phosphorylation of Dazl. MK2 plays an important role in somatic cells in the mitotic G2/M checkpoint in response to DNA damage through direct phosphorylation of Cdc25 (Manke et al., 2005). Our results indicate that MK2 plays a similar, albeit indirect, role in meiosis through phosphorylation of Dazl, arresting the meiotic cell cycle due to reduced translation of Twine.

The primary phenotype of MK2-knockout mice is increased resistance to lipopolysaccharide-induced endotoxic shock due to a dampened inflammatory response (Kotlyarov et al., 1999). Our data suggest that in addition to being more resistant to inflammation, MK2-knockout male mice may also display an increased tolerance to heat-induced germ cell apoptosis. Although this remains to be tested, it opens up the possibility that MK2 kinase inhibitors may find utility in the treatment of male infertility.

MATERIALS AND METHODS

Antibodies and reagents

Anti-Dazl (E6), anti–c-Myc (9E10), anti-GST (Z-5), anti–β-tubulin (H-235), anti–glyceraldehyde-3-phosphate dehydrogenase (6C5), anti-Vasa (d-260), and anti-PABP (10E10) antibodies were from Santa Cruz Biotechnology (Dallas, TX). Anti-MK2, anti–phospho-MK2 (T222), anti–phospho-MK2 (T334), anti–phospho-p38 (T180/Y182), anti–Hsp27, and anti–phospho–Hsp27 (S82) antibodies were from Cell Signaling Technology (Danvers, MA). Anti-HA antibody (16B12) was from Covance (Princeton, NJ). Anti–thiophosphate ester antibody (51-8) and p-nitrobenzyl mesylate (PNBM) were from Abcam (Cambridge, MA). ATP, ATPγS, 4′,6-diamidino-2-phenylindole, and sodium arsenite were from Sigma-Aldrich (St. Louis, MO). The p38 MAPK inhibitor SB203580 was from Selleckchem (Houston, TX), and MK2 inhibitor III was from Cayman Chemical (Ann Arbor, MI).

Cell culture

HEK293 and HeLa cells were grown in DMEM supplemented with 10% fetal bovine serum (FBS) and penicillin/streptomycin. SUM149 cells were grown in Ham’s F12 medium supplemented with 5% FBS, penicillin/streptomycin, insulin (5 μg/ml), and hydrocortisone (1 μg/ml). Cells were grown at 37°C in 5% CO2. Lipofectamine 2000 (Invitrogen, Carlsbad, CA) was used for transfection according to the manufacturer’s instructions. To induce oxidative stress, cells were stimulated with 0.5 mM sodium arsenite for 20 min at 37°C. To prepare lysates for immunoblotting or immunoprecipitation, cells were lysed by scraping in ice-cold lysis buffer (20 mM Tris, pH 8.0, 200 mM NaCl, and 1% Triton X-100 with protease inhibitors [Roche, Indianapolis, IN] and phosphatase inhibitors [1 mM sodium orthovanadate, 10 mM sodium fluoride, and 1 mM β-glycerophosphate]).

Generation of MK2-knockdown cell line

Retroviral shRNA constructs targeting human MK2 and a scrambled control shRNA construct were purchased from OriGene (Rockville, MD). Stably transfected SUM149 cells were selected with puromycin
(1 μg/ml) were maintained in medium containing puromycin (0.5 μg/ml). Knockdown of MK2 was confirmed by Western blotting.

Plasmids
Human MK2 was PCR amplified as an EcoRI–NotI fragment from a cDNA clone (OriGene) and cloned into the corresponding sites of pCMV-myc (Clontech, Mountain View, CA). A plasmid encoding dominant-negative p38 (FLAG-p38 alpha [AF]) was provided by Roger Davis (University of Massachusetts Medical School, Worcester, MA; plasmid #20352; Addgene). A construct encoding constitutively active MK2 (pCDNA3-myc-MK2-EE) was provided by Matthias Gaestel (Hannover Medical School, Germany). A plasmid encoding human PABP was purchased from OriGene. Human Dazl was PCR amplified from a cDNA clone (OriGene) as an EcoRI–BglII fragment and cloned into the corresponding sites of pCMV-myc (Clontech) and pCMV-HA (Clontech). The S65A, S65D, and R115G mutations were introduced by PCR-mediated site-directed mutagenesis using the QuikChange Kit (Agilent, Wilmington, DE). Oligonucleotides were purchased from Integrated DNA Technologies (Coralville, IA). Primer sequences are available upon request. All constructs were fully sequenced.

Protein microarrays and data analysis
The Snapshot Proteomics screen was conducted by AVMBioMed (Malvern, PA). Protein microarrays (consisting of >20,000 human proteins) were blocked for 1 h at room temperature in phosphate-buffered saline (PBS) containing 0.05% Tween-20, 20 mM reduced glutathione, 1 mM dithiothreitol (DTT), 1% bovine serum albumin, and 25% glycerol. Duplicate arrays were incubated for 90 min at room temperature using the two cell lysates (control shRNA and MK2 shRNA) at 2 mg/ml total protein concentration, followed by three washes with 1× PBS. Arrays were then incubated for 1 h at room temperature with rabbit anti–phosphoserine (Cell Signaling Technology) and rabbit
anti-phosphothreonine (Abcam) antibodies per the manufacturer’s instructions for immunoblotting. Arrays were washed three more times, followed by a 1-h incubation with Alexa 488-conjugated anti-rabbit secondary antibody. This was followed by two washes with PBS/Tween-20, PBS, and then two washes with water, followed by centrifugal drying (1000 rpm for 5 min at room temperature) and scanning (GenePix 4100A; Molecular Devices) of the arrays.

Data were obtained from 21,685 proteins present on the microarray. Microarray images were gridded and quantitated using GenePix Pro (version 7) software. Median intensities (features and local backgrounds) were used, and signal-to-noise ratio (SNR) was calculated. Duplicate features representing identical proteins were summarized by average (avg M) and SD. Magnitude change was calculated as log(SNRcontrol RNAi) − log(SNRMK2 RNAi). These values were then Loess transformed by print tip and location to remove technical sources of error (Smyth and Speed, 2003), resulting in the final estimate of magnitude change (M value). To assess statistical significance, a paired, two-tailed Student’s t test was used to calculate a p value for each estimate using the null hypothesis that avg M = 0. A threshold of 95% confidence (p < 0.05) and a cutoff of M > 1.0 were applied to filter the data.

Fly stocks and transgenic fly lines
Stocks were maintained and all crosses performed at 25°C on molas-based food. bol flies were kindly provided by Steve Wasseman (University of California, San Diego, La Jolla, CA). Human DAZL or DAZL-S65D was PCR amplified using attB-modified primers and cloned into pDONR-Zeo (Invitrogen). The cloned inserts were transfected into pTMW using Gateway cloning technology to create N-terminal myc-tagged constructs for Drosophila embryo micro-injection. Transgenic flies were generated by BestGene (Chino Hills, CA). UAS-DAZL maps to the third chromosome, and UAS-DAZL-S65D maps to the second chromosome.

Protein expression and purification
BL21(DE3) cells (Novagen, Billerica, MA) were transformed with plasmids encoding GST, GST-Dazl, GST-Dazl-S65A, GST-Dazl-RRM-WT, GST-Dazl-RRM-S65D, or GST-Dazl-RRM-R115G. Protein expression was induced at 37°C for 2 h with 1 mM isopropyl-β-D-thiogalactoside (IPTG). After centrifugation, the pellet was resuspended in 15 ml of lysis buffer (20 mM Tris, pH 7.4, 150 mM NaCl, 1 mM DTT, and 1 mM EDTA plus protease inhibitors (Roche)). The lysate was sonicated on ice and centrifuged at 20,000 × g for 20 min at 4°C. The supernatant was added to 0.5 ml (packed volume) of glutathione agarose beads (Gold Biotechnology, St. Louis, MO) and incubated at 4°C for 2 h with end-over-end tumbling. The slurry was transferred to a column and washed extensively with wash buffer. GST-fusion proteins were eluted with excess glutathione in lysis buffer (pH 7.4), and appropriate fractions were pooled. The protein was dialyzed overnight at 4°C into dialysis buffer (20 mM Tris, pH 7.4, 150 mM NaCl, 1 mM DTT). Glycerol was added to a final concentration of 10%, and the proteins were snap-frozen and stored at −80°C.

In vitro kinase assays
Recombinant GST-tagged MK2 (residues 40–400; R&D Systems, Minneapolis, MN) was incubated with 20 μg of recombinant purified Dazl protein in kinase buffer (50 mM HEPES, pH 7.5, 0.65 mM MgCl2, 0.65 mM MnCl2, 12.5 mM NaCl) with 500 μM ATP or ATPγS, as indicated. Kinase reactions were incubated at 30°C for 30 min. When appropriate, PNB (dissolved in dimethyl sulfoxide (DMSO)) was added to a final concentration of 2.5 mM, and reactions were incubated at room temperature for 1 h, followed by the addition of 6× boiling sample buffer to stop the reaction. Reactions were then resolved by SDS-PAGE, followed by immunoblotting with the indicated antibodies.

Generation of anti–phospho-S65-dazl antibody
A custom phosphospecific antibody was generated by GenScript (Piscataway, NJ). New Zealand white rabbits were injected with a phosphopeptide (ARYGPSVEKVTDC) conjugated to KLH. Sera were tested for sensitivity and specificity by enzyme-linked immunosorbent assay. The antibody was affinity purified against the phosphorylated peptide and negatively adsorbed against the nonphosphorylated peptide.

SDS–PAGE and Western blotting
Proteins were resolved on 12.5% polyacrylamide gels and transferred to nitrocellulose membranes (GE Amersham) using the Pierce G2 Fast Blotter (Thermo Scientific, Waltham, MA). Membranes were blocked in 5% nonfat milk in Tris-buffered saline (TBS), followed by overnight incubation with the indicated primary antibody in the same buffer. Membranes were washed the next day in TBS plus 0.5% Tween (TBS-T), incubated for 1 h with the appropriate horse-radish peroxidase–conjugated secondary antibody, washed again in TBS-T, and processed for signal detection using enhanced chemiluminescence (Santa Cruz Biotechnology). For detection of proteins from the same reaction/lysate that migrate at similar molecular weights (i.e., phosphorylated and nonphosphorylated forms of the same protein), samples were split and run on separate gels.

Surface plasmon resonance
For protein expression for use in SPR, all plasmid DNA constructs were transformed into BL21(DE3) cells (Life Technologies). Cells were grown in Terrific Broth (RPI, Mount Prospect, IL) at 37°C in 2- to 8-l volumes using a LEX bioreactor (EpiphYTE, Toronto, Canada), induced using 100 μM IPTG (CellGro, Manassas, VA) at 15°C, and harvested 12 h later by centrifugation at 4000 × g for 20 min. Pellets were stored at −80°C until purification. Bacterial cell pellets from 2-l culture volumes were thawed in warm water and then resuspended using binding buffer (50 mM Tris, pH 8.0, 500 mM NaCl) plus a 3:1 ratio of B-PerY-Per (Thermo Fisher), DNase 1, lysozyme, and protease inhibitor P2714 (all from Sigma-Aldrich). Cells were sonicated (frequency 60% maximum, 30 s on/30 s off, 5-min sonication time), and total lysate was mixed with 1–2 ml of His-Link resin (Promega, Madison, WI) and then incubated with rocking for 1 h at 4°C. His-Link resin was cleaned in batch mode by extensive washing with cold binding buffer and transferred into gravity columns (Bio-Rad, Hercules, CA) for final washes (with wash buffer: binding buffer plus 10 mM imidazole, pH 8.0) and elution (with elution buffer: binding buffer plus 250 mM imidazole, pH 8.0, and 10% glycerol). Size exclusion chromatography for all proteins was accomplished using a Sephadex 200 column (GE Healthcare) on an NGC Chromatography System (Bio-Rad) using gel filtration buffer (binding buffer plus 5 mM β-mercaptoethanol and 1 mM EDTA). Fractions were analyzed by SDS–PAGE and pooled. If needed, protein was concentrated using Vivaspin-20 concentrators with a 5000–molecular weight cut-off (GE Healthcare).

Experiments were performed using the ProteOn XPR36 SPR array system (Bio-Rad). ProteOn GLC sensor chips were preconditioned with two short pulses each (10 s) of 50 mM NaOH, 100 mM HCl, and 0.5% SDS. Then the system was equilibrated with PBS-T buffer (20 mM Na phosphate, 150 mM NaCl, and 0.1% polysorbate 20, pH 7.4). Individual ligand flow channels were activated for 5 min
at 25°C with a mixture of 1-ethyl-3-(3-dimethylaminopropyl carbodiimide hydrochloride) (0.2 M) and sulfo-N-hydroxysuccinimide (0.05 M). Immediately after chip activation, NeutrAvidin (Thermo-Fisher) at a concentration of 100 µg/ml in 10 mM sodium acetate, pH 5.0, was injected across ligand flow channels for 5 min at a flow rate of 30 µl min⁻¹. Excess active ester groups on the sensor surface were capped by a 5-min injection of 1 M ethanolamine HCl (pH 8.5). This resulted in the coupling of NeutrAvidin at a density of 9000 response units (an arbitrary unit that corresponds to 1 pg/mm²). The SD in the immobilization level from the six spots within each channel was <4%. Synthetic 3’-biotinylated RNA oligonucleotides (IDT) were then affinity purified and used as ligand for the Dazl-RRM protein–binding experiments.

Dazl-RRM proteins at the indicated concentrations in RNA-protein binding buffer (20 mM Tris, pH 7.5, 100 mM NaCl, 1 mM MgCl₂, 0.005% Tween) were injected over the control (no oligonucleotide) and oligonucleotide surfaces at a flow rate of 100 µl/min for a 2-min association phase, followed by a 4-min dissociation phase at 25°C using the “one-shot” functionality of the ProteOn (Bravman et al., 2006). Specific regeneration of the surfaces between injections was not needed due to the nature of the interaction. Data were analyzed using ProteOn Manager Software, version 3.0 (Bio-Rad). The responses of a buffer injection and responses from the reference flow cell were subtracted to account for nonspecific binding. Data were subjected to equilibrium analysis, plotting the response at equilibrium versus concentration and fitting to a steady-state model to obtain the equilibrium dissociation constant (Kd).

**Dual luciferase assay**

The 3’ UTR of SYCP3 was PCR amplified from mouse genomic DNA as an Xhol–Notl fragment and cloned into the corresponding sites of psiCHECK-2 (Promega). HeLa cells were seeded in a 96-well plate and grown overnight to ~65% confluence. Cells were cotransfected with plasmids encoding myc-tagged Dazl, Dazl-S65D, or Dazl-R115G and psiCHECK-2/SYCP3-3’UTR or transfected with just psiCHECK-2/SYCP3-3’UTR alone. The next day, cells were lysed according to the protocol described in the dual luciferase reporter assay kit (Promega). Cell lysates were transferred to wide-bottom 96-well plates that were read on a GLOMAX-96 microplate luminometer. Firefly luciferase values were normalized to Renilla luciferase values for each individual condition (performed in triplicate). These values were averaged, and the Dazl-S65D and Dazl-R115G values were normalized to the values obtained from Dazl-WT. Statistical analysis was performed using a Student’s t test to obtain a p value.

**Hatch-rate analysis**

Two males of the indicated genotype were crossed to five virgin w¹¹¹ females for 2 d. Flies were then transferred to a collection cage attached to a grape juice agar plate for overnight collection of deposited eggs. The hatch rate was determined by dividing the number of hatched embryos after 24 h by the total number of embryos deposited at 14 h.

**Testes dissection and microscopy**

Testes from 1-d-old male flies were dissected in PBS and prepared for microscopy as described (Sitaram et al., 2014). Live testes were imaged by phase contrast microscopy using an EVOS FL Auto microscopy (Life Technologies) equipped with a 10x objective. For immunofluorescence microscopy using HeLa cells, cells were seeded on glass coverslips and transfected the next day with the indicated constructs for 24 h. Cells were then fixed in 4% formaldehyde and processed for immunofluorescence microscopy. Cells were imaged using an EVOS FL Auto microscope equipped with a 40x objective.

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**Phosphorylation of Dazl by MK2**

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