Loss of the *Drosophila melanogaster* DEAD box protein Ddx1 leads to reduced size and aberrant gametogenesis

Devon R. Germain\textsuperscript{a}, Lei Li\textsuperscript{a}, Matthew R. Hildebrandt\textsuperscript{a}, Andrew J. Simmonds\textsuperscript{b}, Sarah C. Hughes\textsuperscript{b,c}, Roseline Godbout\textsuperscript{a,}\textsuperscript{*}

\textsuperscript{a} Departments of Oncology University of Alberta, Cross Cancer Institute, 11560 University Avenue, Edmonton, Alberta, T6G 1Z2, Canada
\textsuperscript{b} Cell Biology University of Alberta, Cross Cancer Institute, 11560 University Avenue, Edmonton, Alberta, T6G 1Z2, Canada
\textsuperscript{c} Medical Genetics University of Alberta, Cross Cancer Institute, 11560 University Avenue, Edmonton, Alberta, T6G 1Z2, Canada

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**A B S T R A C T**

Mammalian DDX1 has been implicated in RNA trafficking, DNA double-strand break repair and RNA processing; however, little is known about its role during animal development. Here, we report phenotypes associated with a null *Ddx1* (*Ddx1\textsuperscript{fsy}*) mutation generated in *Drosophila melanogaster*. *Ddx1* null flies are viable but significantly smaller than control and *Ddx1* heterozygous flies. Female *Ddx1* null flies have reduced fertility with egg chambers undergoing autophagy, whereas males are sterile due to disrupted spermatogenesis. Comparative RNA sequencing of control and *Ddx1* null third instars identified several transcripts affected by *Ddx1* inactivation. One of these, *Sirup* mRNA, was previously shown to be overexpressed under starvation conditions and implicated in mitochondrial function. We demonstrate that *Sirup* is a direct binding target of Ddx1 and that *Sirup* mRNA is differentially spliced in the presence or absence of *Ddx1*. Combining *Ddx1* null mutation with *Sirup* dsRNA-mediated knock-down causes epistatic lethality not observed in either single mutant. Our data suggest a role for *Drosophila* *Ddx1* in stress-induced regulation of splicing.

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1. Introduction

DEAD box proteins are a family of RNA helicases implicated in virtually every aspect of RNA metabolism (Cordin et al., 2006). These proteins are characterized by 12 conserved motifs, including the D(Asp)-E(Glu)-A(Ala)-D(Asp) motif for which they are named. DEAD box proteins function by modifying RNA secondary structure in an ATP-dependent manner (Linder and Fuller-Pace, 2013), and play a role in RNA trafficking (Linder and Stutz, 2001; Montpetit et al., 2012). There are > 30 DEAD box genes in the *Drosophila melanogaster* genome (2014).

Several *Drosophila* DEAD box genes are known to play a role in early development and gametogenesis. For example, the *vasa* gene encodes a multifunctional DEAD box protein that localizes to the posterior pole in oocytes and is required for completion of oogenesis (Lasko, 2013). *Vasa* has also been implicated in chromatin condensation and generation of small non-coding RNAs (Pek and Kai, 2011; Zhang et al., 2012). *Belle*, which is closely related to *Vasa*, is essential for larval development and required for both male and female fertility (Johnstone et al., 2005). Mutation of *pitchoune*, encoding another DEAD box helicase, results in developmental arrest during the first instar stage. *Pitchoune* has been implicated in regulating cell growth and proliferation (Zaffran et al., 1998).

Only one *Ddx1* mutation has been previously described in *Drosophila melanogaster* (Zinsmaier et al., 1994). This mutation was deemed to be recessive lethal; however, the nature of the mutation was not determined and the mutant line is no longer available. Rafti et al. described the expression of *Ddx1* in *Drosophila* in 1996, and reported elevated levels in early embryos, and expression throughout development (Rafti et al., 1996). More recently, publicly available large scale studies using gene expression microarrays and RNA deep sequencing analysis have revealed widespread expression of *Ddx1* in all tissues and cell lines tested to date (St Pierre et al., 2014). These screens show elevated *Ddx1* levels in the nervous system, testes and ovaries with the highest levels observed at early embryonic stages, as previously described (Rafti et al., 1996).

Human DDX1, which is 77% similar to *Drosophila* *Ddx1*, is amplified and overexpressed in a subset of MYCN-amplified neuroblastoma and retinoblastoma cell lines and tumors (Godbout et al., 1998; Godbout and Squire, 1993; Manohar et al., 1995; Squire et al., 1995). *Ddx1* is also a prognostic marker in breast cancer (Balko and Arteaga, 2011; Germain et al., 2011), and plays a role in...
testicular tumorigenesis (Tanaka et al., 2009). Large scale screens for disease associated genes have identified DD1X as a potential gene of interest in cervical cancer and chronic obstructive pulmonary disease (Johanneson et al., 2014; Smolonska et al., 2014). As well, DD1X is significantly down-regulated in Down syndrome fetal brains (Kircher et al., 2002). Clues to DD1X's role in these diseases come from in vitro analysis. For example, DD1X was identified in a subset of RNA transport granules involved in the subcellular localization of RNA molecules in neuronal axons (Kanai et al., 2004). DD1X’s role in RNA trafficking is not limited to endogenously coded genes as HIV replication requires DD1X for efficient export of unspliced viral genomic RNA from the nucleus to the cytoplasm (Edgcomb et al., 2012; Fang et al., 2005; Robertson-Anderson et al., 2011). This effect is mediated through an interaction between DD1X and two virally encoded proteins, Rev and Tat, that are essential for RNA export (Lin et al., 2014). DD1X is also required for efficient replication of coronavirus (Wu et al., 2014; Xu et al., 2010), and has been shown to transactivate hepatitis C and JC viral genes (Sunden et al., 2007a; Sunden et al., 2007b; Tingting et al., 2006).

Under normal conditions, DD1X is mainly found in the nucleus where it forms granules that co-localize with (or reside adjacent to) nuclear organelles associated with mRNA processing such as cleavage bodies, gems and Cajal bodies (Bleoo et al., 2001; Li et al., 2006). When cells are treated with ionizing radiation, DD1X is recruited to a subset of DNA double-strand breaks (Li et al., 2008). Biochemical analysis has shown that DD1X can unwind RNA/RNA and RNA/DNA duplexes in vitro in an ADP-dependent manner and can efficiently digest single-stranded RNA (Li et al., 2008). Finally, DD1X has also been implicated as a mRNA splicing factor (Popow et al., 2014).

While it is clear that DD1X is a multifunctional protein, we only have a limited understanding of its biological role during development. Therefore, in order to gain insight into DD1X’s role during development, we generated a new DD1X null mutant Drosophila line. Here, we report that DD1X null flies are viable, with reduced fertility and body size. DD1Xnull flies also display aberrant gametogenesis in both testes and ovaries. We also describe a direct interaction between Sirup mRNA, previously described as up-regulated during starvation conditions and implicated in miotochondrial function (Erdi et al., 2012; Van Vranken et al., 2014), and DD1X protein. Finally, Sirup RNA is differentially spliced in control versus DD1X null flies, and an epistatic lethal effect is observed in DD1X+/+/Sirup dsRNA-mediated knock-down flies.

2. Materials and Methods

2.1. Drosophila Stocks

All crosses were performed at 25°C on standard Bloomington recipe media. Fly stocks were obtained from the Vienna Drosophila RNAi Center (VDRC) (Vienna, Austria) and the Bloomington Drosophila Stock Center (BDSC) (Indiana University, USA). The following fly stocks were used:

- w1118, Control line
- yw3; Ly/TM3, Sb – Balancer for potential mutant allele
- yw67C2; P[EPgy2]EY12792 – P-element upstream of DD1X
- w; Dr/TM3, P[ry]{Δ2-3}1998, BSDC #1610, Individual white-eyed F1 single female virgin w/+; ΔP[EPgy2]EY12792/TM3, P[ry]{Δ2-3}1998 flies were mated to yw/+; Ly/TM3, Sb males. Individual F2 w/+; ΔP[EPgy2]EY12792/TM3, Sb flies were then again mated to yw/+; Ly/TM3, Sb flies, and w/+; ΔP[EPgy2]EY12792/TM3, Sb female and male progeny from each cross were further crossed to establish balanced lines.

2.2. Generation of potential Ddx1 mutant alleles by P-element excision

Potential mutations of the Ddx1 locus were generated using BDSC stock #21389 (yw flw67C2; P[EPgy2]EY12792), that contains a P-element inserted 43 bp upstream of the Ddx1 transcriptional start site. P[EPgy2]EY12792 was excised by crossing virgin P[EPgy2]EY12792 females with Δ2-3 transposase expressing males (w; Dr/TM3, P[ry]{Δ2-3}1998, BSDC #1610). Individual white-eyed F1 single female virgin w/+; ΔP[EPgy2]EY12792/TM3, P[ry]{Δ2-3}1998 flies were mated to yw/+; Ly/TM3, Sb males. Individual F2 w/+; ΔP[EPgy2]EY12792/TM3, Sb flies were then again mated to yw/+; Ly/TM3, Sb flies, and w/+; ΔP[EPgy2]EY12792/TM3, Sb female and male progeny from each cross were further crossed to establish balanced lines.

2.3. Characterization of potential mutant lines

Individual fly lines displaying white eye color, indicating removal of P[EPgy2]EY12792 (which carries a w+ transgene), were analyzed for deletion by sequential PCR reactions using staggered primers. One line (AX) was identified as containing an ~1.7 kb deletion. Subsequent genomic sequencing revealed that a 1733 bp region (3L: 22,263,326 to 22,265,059) spanning the P-element insertion site and most of the Ddx1 gene locus had been removed and replaced with the 15 bp sequence 5′-CATGATGAAATAAACAG-3′. This 15 bp sequence does not correspond to any part of Ddx1 or P[EPgy2]EY12792. This allele was designated Ddx1AX. The following primers were used for PCR amplification: 5′-CCAAGACCGTTAGTC-3′ (forward primer ~400 bp upstream of Ddx1 transcription start site), 5′-ATGAGGTGTGGCCAGCG-3′ (forward primer ~500 bp downstream of Ddx1 transcription start site), 5′-AGCTGTGGAATGGACG-3′ (reverse primer ~400 bp downstream of Ddx1 transcription start site), 5′-ACACCTGCAGACGG-3′ (reverse primer ~1.4 kb downstream of Ddx1 transcription start site), 5′-GACCTCCGACTTCTAC-3′ (reverse primer located in Ddx1 3′ UTR). The following primer was used for genomic DNA sequencing: 5′-CTCTAAAGTCAAGTACAC-3′ (forward primer ~200 bp upstream of Ddx1 transcription start site) (Integrated DNA Technologies, Coralville, Iowa).

2.4. Western blot analysis and antibodies

Cell lysates were prepared from adult flies or larvae by grinding with a pestle in lysis buffer (1% sodium deoxycholate, 1% Triton-X-100, 0.2% SDS, 150 mM NaCl, 50 mM Tris-HCl pH 7.4 and 1X Complete protease inhibitors (Roche, Mississauga, Canada). Samples that were analyzed for phospho-S6k were prepared in lysis buffer supplemented with 1X PhosStop (Roche). Lysates were electrophoresed in 10% SDS-polyacrylamide gels and transferred to nitrocellulose membranes. The following antibodies were used: rabbit anti-pS6k (1:1000) (#9209, Cell Signaling Technology, New England Biolabs, Whitby Canada), rabbit anti-DD1X (1:500) (custom polyclonal antibody, antigen CQNLKRTGSYEDHV, GenScript, Piscataway, USA) and mouse anti-β-tubulin (1:2000) (DSHB, E7). The signal was detected using secondary anti-rabbit-HRP and anti-mouse-HRP antibodies (Jackson ImmunoResearch, West Grove, USA). The β-tubulin antibody developed by Michael Klymowski was obtained from the Developmental Studies Hybridoma Bank, created by the NICHD of the NIH and maintained at the University of Iowa, Department of Biology, Iowa City, IA 52242. Proteins were detected using the Immobilon (EMD Millipore, Darmstadt, Germany).
2.5. RNA purification and RT-PCR

RNA was purified using an RNeasy Plus Universal Mini Kit (Qiagen, Toronto, Canada) as per the manufacturer's directions. Briefly, whole flies or larvae were crushed in QiAzo1 reagent using plastic pestles and purified RNA isolated using a mini column. Reverse transcription was carried out using SuperScript II reverse transcriptase (Life Technologies, Burlington, Canada) and oligo dT primers as per the manufacturer's directions. The following primers were used for RT-PCR analysis: Sirup, forward primer 5'-CCTGCGAGATTGCAAATTC-3', reverse primer 5'-AGTGGTCTCTTCTCCTGG-TAGC-5'; Sirup splice specific transcript, forward primer 5'-CAGAAAAGGGCAAACAAAGTA-3' (asterisk indicates splice junction site), reverse primer 5'-GAATTCCTTATAAGTTCCTGGC-3'; actin, forward primer 5'-AAATCATAGAACACAAA-3', reverse primer 5'-GAAGTAGACATGTTAGTACG-3'. The forward primer for Sirup splice-specific product amplification consisted of the 15 nt sequence upstream of the splice junction followed by the 4 nt sequence downstream of the splice junction.

2.6. Viability, fertility, size and developmental assays and larval collections

For viability and fertility assays, single virgin females were mated with two males and left to lay eggs in standard culture tubes. After ten days, the parental flies were removed and individual pupae were counted and moved to a new tube daily to ensure no cross-generational contamination. For crowded conditions, twenty females were mated to twenty males in a single tube. For size analysis, newly eclosed adults or pupae were genotyped and photographed using an Olympus SZX12 fluorescence dissecting microscope coupled to a Canon Powershot G2 color camera. Whole length, for pupae, or thorax length, for adults, was measured using Photoshop coupled to a Canon Powershot G2 color camera. Whole length was measured using a ruler with standard parameters. Differential gene expression was calculated using the NIH and maintained at the University of Iowa, Department of Biology, Iowa City, IA 5224.

Confocal images were captured on a Zeiss LSM 710 confocal laser scanning microscope with a plan-Apochromat 63x (NA 1.4) oil immersion lens, a plan-Apochromat 40x (NA 1.3) oil immersion lens, or a plan-Apochromat 10x (NA 0.45) lens, and Zen software v7.04.287. An Olympus SXZ12 microscope was used to photograph adult flies and gonads. Exported images were saved as TIFF files and measurements made using Photoshop.

2.8. Northern blot analysis

RNA samples were isolated from wandering 3rd instars as described above. For each sample, 5 μg of total RNA were resolved in a 10% denaturing urea polyacrylamide gel. A small RNA ladder (NEB, Whitby, Canada) was used for size determination. RNA was then transferred to Hybond-N+ membranes (GE Healthcare, Mississauga, Canada) and baked for 1 hour at 80 °C. DNA probes were generated by 32P end-labeling of single-stranded oligonucleotides. The following probes were used: tRNA5'−CCAATGGGCAAACAAAGTA−3', tRNA5'−TTGCTAAGGCAGGCGTCTCCTACTGACGTAACCC−3', 5s RNA 5'−CAGCGTCACGGGTTAGTACG−3', and 5s RNA 5'−TCATCGGGCTTACGTAACCAAGCAGCAGAC−3'. Probe hybridization was carried out in 5X SSC, 50 mM Na2HPO4 pH 6.5, 0.5X Denhardt's solution and 0.25 mg/mL salmon sperm DNA at 42 °C. The blot was then washed in 2X SSC, 0.1% SDS, followed by 0.1X SSC, 0.1% SDS. The signal was visualized using X-ray film. Membranes were initially hybridized with radioactively-labeled tRNA5' and 5s RNA probes. The label was then removed using stripping buffer and the blot exposed to film to ensure that the signal was sufficiently reduced to allow re-probing. The membrane was then re-hybridized with radioactively-labeled tRNA5' probe.

2.9. RNA deep sequencing

RNA was isolated from three independent preparations of wandering 3rd instars for both Ddx1 null and control genotypes. RNA libraries were prepared by first removing rRNAs using a Ribozero™ tRNA Removal Kit (Human/Mouse/Rat) (Epitope, Madison, USA) followed by TruSeq Stranded Total RNA Sample Prep Kit (Illumina, San Diego, USA). Paired-end 100 nt RNA sequencing was performed on an Illumina HiSeq 2000 platform and processed using Casava V1.8.2 by the UBC Biodiversity Research Centre NextGen Sequencing Facility. Alignment, splice site identification was carried out using Bowtie v2.1.0 and Tophat v2.0.13 with standard parameters. Differential gene expression was calculated using Cufflinks v2.1.1 using standard parameters with a P-value of <0.01 considered significant.

2.10. RNA co-immunoprecipitations

Whole cell lysates were prepared from S2 cells by resuspending cell pellets in 50 mM Tris–HCl pH 7.5, 150 mM NaCl, 0.5% sodium deoxycholate, 1% NP-40, and 1X Complete (Roche) protease inhibitors. Four hundred micrograms of lysate were first cleared with Protein A agarose beads (GE Healthcare), followed by incubation with 5 μl of rabbit anti-Ddx1 antibody or rabbit IgG for 2 hours at 4°C. Protein A agarose beads were then added and incubation continued for 1 hour at 4°C. Co-immunoprecipitates were washed three times in lysis buffer and extracted with water-saturated phenol. An aliquot taken from the co-immunoprecipitates was saved for western blot analysis to check Ddx1 immunoprecipitation efficiency. Co-immunoprecipitated RNA was resuspended in RNase-free water and stored at -80°C until use. Reverse transcription was carried out as described above, using Sirup-specific (5'-AGTGGTCTCTTCTCCTGGTACG-3') or Ddx1-specific (5'-TCATCGGGCAGCGTACACG-3') reverse primers with the immunoprecipitated RNA.
serving as template. Following reverse transcription, PCR amplification was carried out using Sirup specific (forward 5'-CCTCGGAGATTGCAATTACAG-3', reverse 5'-AGTGTTCTTCTTCGGTAGC-5') or Ddx1 specific (forward 5'-GCATTGGTAGGGAAG-3', reverse 5'-TCATCGGAGGCACACG-3') primers.

2.1. Sirup knock-down in Ddx1 modified flies

yw*; P(Act5C-GAL4)25FO1/CyO; Ddx1AX/TM3, Sb virgin females were crossed to w1118; P(GD14644)v36437; Ddx1AX/TM3, P(ActGFP) JMR2, Ser1 males and allowed to lay eggs for 3 days. Adults were scored for Ddx1 status by the presence of Sb or Ser alleles, and Sirup knock-down was scored based on the absence of CyO. Chi square analysis was performed to determine the significance of observed outcomes compared with expected genotype distribution.

3. RESULTS

3.1. Ddx1 null flies are viable

Imprecise excision of a P-element located immediately upstream of Ddx1 (yw67C23, P[Epgy2]EY12792) was used to screen for novel Ddx1 mutations. From this screen, we identified a new null Ddx1 allele with a 1733 bp deletion encompassing the majority of the Ddx1 open reading frame. This allele was designated Ddx1AX (Fig. 1A). In contrast to a previously reported Ddx1 mutation...

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**Fig. 1. Ddx1 null flies are viable but show reduced fertility.** (A) The location of the Ddx1AX deletion. (B) Western blot analysis shows no detectable signal for Ddx1 protein in Ddx1AX/AX adult flies. (C) Survival of homozygous adult flies generated from Ddx1 heterozygote crosses. As the Ddx1AX mutation is carried over a recessive lethal balancer chromosome, the expected percentage of homozygous progeny is 33%. At low density, homozygous flies were generated at the expected rate. At medium density and high density, a significant reduction in the number of homozygous flies was observed; n=45 adults (low density), 1165 adults (medium density) and 499 adults (high density). (D) Progeny generated from single virgin females mated with two males (genotypes are indicated) and allowed to lay eggs for 10 days. Pupae were removed and counted daily. Homozygous mutant flies generated very low or no progeny. Heterozygous flies generated progeny at the expected rate. n ≥ 20 crosses for all samples. Error bars indicate s.e.m.
(Ddx1<sup>w118</sup>) that showed early embryonic lethality (Zinsmaier et al., 1994), Ddx1<sup>AX/AX</sup> flies reached adulthood. Western blot analysis of control (w<sup>1118</sup>) and Ddx1<sup>AX/AX</sup> adult flies showed a complete absence of Ddx1 in homozygous mutants (Fig. 1B). Although a small portion of the 3’ Ddx1 open reading frame is retained in Ddx1<sup>AX/AX</sup> flies, no novel bands were detected using an anti-Ddx1 antibody generated against the C-terminus of Ddx1. These results suggest that there is either no translation of the retained 3’-end of Ddx1 or the resulting product is unstable.

Although Ddx1<sup>AX/AX</sup> flies were viable, we noticed that they were consistently outcompeted under high density culturing conditions. When single eggs from heterozygote crosses were raised in isolation, we observed the expected 2:1 ratio of heterozygous to homozygous mutant progeny (heterozygous mutants were maintained over a recessive lethal balancer chromosome [TM3, P {ActGFP}JMR2, Ser]<sup>1</sup>, so no homozygous non-mutant Ddx1 progeny were generated). When a single female was allowed to lay eggs for 10 days in a standard collection tube, the ratio dropped to approximately 3:1. When 20 females were allowed to lay eggs in a standard collection tube for 10 days, the ratio of heterozygous to homozygous mutant was 13:1 (Fig. 1C).

3.2. Ddx1 null flies have reduced fertility, size and delayed development

Heterozygous y<sup>1</sup>w<sup>52</sup>; Ddx1<sup>AX/TM3</sup> Ser GFP flies generated the expected number of progeny as compared to control flies (Fig. 1D). Both male and female Ddx1<sup>AX/AX</sup> flies were sterile. To confirm that the observed infertility was due to inactivation of Ddx1, and not a line-specific effect, we out-crossed Ddx1<sup>AX</sup> to a line containing a deficiency that encompasses the Ddx1 locus (Df[3L]ED230). Ddx1<sup>AX</sup>/Df[3L]ED230 males were completely sterile, while females were able to generate a small number of viable progeny that survived to adulthood (Fig. 1D). Heterozygote crosses resulted in approximately 75% as many progeny as control. This was as expected, as 25% of the progeny would be homozygous for the recessive lethal balancer chromosome.

We noted that Ddx1 null progeny generally eclosed later than control and heterozygous animals. To better define the extent of the developmental delay in Ddx1 null flies, we set up heterozygote crosses, with females allowed to lay eggs for a period of 24 hours. Pupae were removed from each collection tube at 24 hour intervals and transferred to a secondary collection tube. Secondary collections were checked for eclosed adults at 24 hour intervals. Time to pupation revealed a non-significant trend (p = 0.069), with heterozygous and homozygous mutant progeny having mean pupation times of 6.07 and 6.38 days, respectively (Fig. 2A, left). A significant delay was observed for time to eclosion (p < 0.001), with heterozygous and homozygous mutant progeny having mean eclosion times of 10.78 and 11.70 days, respectively (Fig. 2A, right).

In order to determine if maternally loaded Ddx1 protein might be compensating for the mutant allele during early development, we examined Ddx1 protein levels in larvae at 24, 48 and 72 hours post egg laying. Significant levels of Ddx1 protein were observed in Ddx1<sup>AX/AX</sup> maternal progeny at 24 and 48 hours post-egg laying, indicating that maternally loaded Ddx1 is still present at these times. Ddx1 was no longer detected at 72 hours.
Ddx1AX/AX larvae at 24 hours, with a weak signal detected as late as 48 hours post egg laying (Fig. 2B). These results suggest a significant amount of maternally deposited protein with a long half-life. This may explain why Ddx1AX/AX flies only display phenotypes at later developmental stages.

Comparison of control, heterozygous and mutant pupal lengths revealed a significant reduction in size in both Ddx1AX/AX and Ddx1AX/Df(3L)ED230 flies (Fig. 3A). Of note, though both null strains were significantly smaller than control animals, the Ddx1AX/Df(3L)ED230 pupae were slightly larger than Ddx1AX/AX. Similar results were observed upon measuring adult length (Fig. 3B). Adult Ddx1 null flies were smaller than control or heterozygous flies, and Ddx1AX/Df(3L)ED230 adults were slightly, though significantly (p < 0.05), larger than Ddx1AX/AX adults.

3.3. Gametogenesis is disrupted in Ddx1 null flies

To investigate the cause of reduced fertility in Ddx1 null flies, we dissected ovaries and testes from adult flies which had been held in isolation for 3 or 10 days following eclosion. At 3 days post-eclosion, heterozygous ovaries appeared essentially identical to...
control, containing many mature eggs. Ovaries from both Ddx1\textsuperscript{AX/AX} and Ddx1\textsuperscript{AX}/Df(3L)ED230 flies were much smaller, with Ddx1\textsuperscript{AX}/Df(3L)ED230 ovaries occasionally containing a small number of mature eggs and Ddx1\textsuperscript{AX/AX} ovaries containing no eggs (Fig. 4A).

As holding flies in isolation on standard media can negatively impact oogenesis (Chapman and Partridge, 1996; Soller et al., 1997), ovaries were isolated from females that were supplemented with yeast paste and allowed to mate. Well-fed mated control and Ddx1 heterozygous females had notably larger ovaries than those kept in isolation, whereas Ddx1\textsuperscript{AX/AX} and Ddx1\textsuperscript{AX}/Df(3L)ED230 ovaries were similar to those from females kept in isolation (Fig. 4B). Similar results were observed with ovaries isolated from adults held in isolation for 10 days post eclosion, with Ddx1\textsuperscript{AX}/Df(3L)ED230 ovaries containing few mature eggs and some Ddx1\textsuperscript{AX/AX} ovaries containing a small number of abnormal eggs that were approximately the size of a mature egg, but lacked dorsal appendages (Fig. 4C).

The reduced size of Ddx1 null flies is likely related to metabolism dysfunction. If this is the case, we would expect to see other phenotypes associated with reduced metabolism. When under metabolic stress or starvation conditions, developing egg chambers undergo autophagy in order to conserve energy (Barth et al., 2011; McCall, 2004). To determine if developing egg chambers in Ddx1 null ovarioles were undergoing autophagy, we carried out immunofluorescence analysis of control and Ddx1 null ovarioles. Early stage egg chambers and germ-line stem cell niches appeared normal, indicating that the early stages of oogenesis are not disrupted in Ddx1 null flies (Fig. 5A). Next, we stained dissected ovarioles with LysoTracker Red, a dye that is commonly used to identify autophagic cells (DeVorkin and Gorski, 2014). In Ddx1 null ovarioles, at both 3 and 10 days post eclosion, we observed mid-stage autophagic egg chambers (Fig. 5B-C, arrows). Autophagic egg chambers were not observed in control and heterozygous ovarioles. As mature egg chambers lacking dorsal appendages were observed in Ddx1 null ovarioles, we imaged stage 8-9 egg chambers of control and Ddx1 null ovarioles for the localization of Gurken, a key factor in dorsal appendage development which localizes to the anterior dorsal corner of mid-stage oocytes (Neuman-Silberberg and Schupbach, 1996; Schupbach, 1987) (Fig. 5D). As expected, control egg chambers displayed normal Gurken localization. While a subset of egg chambers in Ddx1\textsuperscript{AX}/Df(3L)ED230 ovarioles (approximately half) showed the expected Gurken localization pattern, the remaining Ddx1\textsuperscript{AX}/Df(3L)ED230 egg chambers and all egg chambers present in Ddx1\textsuperscript{AX/AX} ovarioles lacked any notable Gurken expression pattern.

Immunofluorescence imaging of testes isolated from adult males revealed aberrant sperm development in Ddx1 null males (Fig. 6A). Early spermatogenesis appeared unaffected in Ddx1 null males, with developing spermatids undergoing nuclear elongation (Fig. 6B). However, elongated spermatids were disrupted and scattered. To determine if mature sperm were being produced, seminal vesicles were isolated and imaged (Fig. 6C). In contrast to control and heterozygous seminal vesicles, which contained mature sperm, we did not observe any mature sperm in the seminal vesicles of homozygous mutant males.

3.4. Ddx1 null flies have reduced ps6k levels, but normal tRNA levels

As both reduced body size and autophagy of developing oocytes are phenotypes associated with reduced metabolism (Barth et al., 2011; Edgar, 2006), we carried out western blot analysis using an antibody that recognizes the phosphorylated form of S6k, a downstream effector molecule of TOR and a common marker of active growth (Montagne et al., 1999). Phospho-S6k is down-regulated under starvation conditions (Hara et al., 1998; Zhang et al., 2000). ps6k levels were notably reduced in Ddx1 null flies at the 3rd instar stage compared to control and heterozygous animals (Fig. 7). Notably, the ps6k signal observed in Ddx1\textsuperscript{AX}/Df(3L)ED230 larvae was stronger than that in Ddx1\textsuperscript{AX/AX} larvae. This is consistent with our previous results demonstrating that the Ddx1\textsuperscript{AX/AX}
Fig. 5. Aberrant development of Ddx1 null egg chambers. Immunofluorescence imaging of ovaries with DAPI and Alexa 546-conjugated phalloidin (A), DAPI and Lyso-Tracker Red (B and C), and DAPI, Alexa 546-conjugated phalloidin and anti-Gurken antibody (D). Developing egg chambers in Ddx1 null ovaries appear normal at early stages (A); however, LysoTracker Red staining reveals egg chambers undergoing autophagy (B and C, white arrows). (D) Immunostaining of stage 8-9 egg chambers with anti-Gurken antibody shows a normal Gurken pattern in control and some Ddx1^{AX}/Ddx1^{AX} egg chambers, with absent Gurken signal in the remaining Ddx1^{AX}/Ddx1^{AX} and all Ddx1^{AX}/Df(3L)ED230 egg chambers.
ED230 phenotype is slightly attenuated compared to Ddx1AX/AX flies.

As human DDX1 has recently been implicated in tRNA splicing, and pS6K levels are associated with RNA Pol III activity (Marshall et al., 2012), we used northern blot analysis to examine tRNA\textsubscript{tyr} (an intron-containing tRNA) and tRNA\textsubscript{ala} (which contains no intron) levels in control, heterozygous and homozygous mutants. Our analysis showed no notable difference in the relative amounts of a mature tRNA that requires splicing (tRNA\textsubscript{tyr}) compared to one that does not require splicing (tRNA\textsubscript{ala}). As well, there was no notable difference in relative tRNA to 5S rRNA levels (Fig. S1).

3.5. Ddx1 null animals display widespread changes in mRNA levels and splicing

DEAD box proteins have been widely implicated in the generation, maturation and degradation of RNA molecules. Therefore, we undertook RNA deep sequencing to determine the effect of knocking-out Ddx1 on the transcriptome. We utilized RNA isolated from control and Ddx1 null 3rd instars, as phenotypes are present at this point and wandering 3rd instars can be easily staged in spite of developmental delays associated with Ddx1 null mutants. We identified 72 significantly down-regulated and 261 significantly
up-regulated transcripts, using a cut-off of \( p < 0.01 \) (Table S1). Of note, we found that RNA is expressed from the portion of the Ddx1\(^{AX} \) allele retained in the genome of our Ddx1 null flies, although RNA levels were significantly reduced compared to that found in control flies. Further analysis of the region directly upstream of the Ddx1\(^{AX} \) allele suggests that transcription is initiated approximately 150 bp upstream of the deletion site. Immediately upstream of this putative transcriptional start site is a variant TATA box sequence (CATAAAG) that may act as a cryptic transcription initiation site in the absence of the normal start site.

We also analyzed transcripts for the presence of differentially spliced variants between Ddx1\(^{AX/AX} \) and control flies (Table S2). We limited our analysis to the small number of transcripts that displayed unique splice variants, with high levels in all three replicates of one genotype and absence in all three replicates of the other genotype (Tables S3-S4). It is important to note that splice site analysis must be considered within the context of total RNA levels, as a gene that is only expressed in one genotype will necessarily have splice junctions that are unique to that genotype. Using these criteria, we identified a number of splice variants uniquely found in either control or Ddx1\(^{AX/AX} \) flies, including Sirup. Gene ontology clustering analysis of genes with modified expression or modified splicing was uninformative.

3.6. Ddx1 interacts with Sirup mRNA

Sirup RNA has previously been shown to be up-regulated in response to starvation conditions (Erdi et al., 2012) and is required for proper mitochondrial function (Van Franken et al., 2014). We observed up-regulation of Sirup (by 3-fold) in Ddx1\(^{AX/AX} \) versus control flies (Table 1). However, a much stronger effect was observed on Sirup splicing, with sequences spanning the 234–315 bp junction found uniquely and at high levels in Ddx1 null flies. Sirup is expressed as a single exon or as an alternatively spliced product characterized by the removal of a small intron located downstream of the start codon (Fig. 8A). Using a primer that spans the junction splice site, we confirmed that Sirup displays alternative splicing in control and null flies, with Ddx1 control larvae and adult flies expressing unspliced Sirup, and Ddx1\(^{AX/AX} \) larvae and adults expressing spliced Sirup (Fig. 8B).

The Sirup splice variant observed in control flies results in a longer transcript (810 nt) that retains the gene’s single intron, but a shorter open reading frame (36 aa) due to a stop codon within the intron. In Ddx1\(^{AX/AX} \)flies, the intron is removed, generating a shorter transcript (662 nt) but a longer open reading frame (118 aa). Intriguingly, this splice-specific regulation may be conserved in humans, as the human Sirup orthologue C6orf57 expresses two splice variants, one encoding a 108 aa protein and the other being non-coding, mirroring what we observe in Drosophila.

As Ddx1 has previously been shown to bind target RNAs, it is possible that Sirup RNA is a direct target of Ddx1. Alternatively, modified splicing of Sirup may be an indirect consequence of Ddx1 mutation. To distinguish between these possibilities, we performed RNA co-immunoprecipitations (co-IPs) using a Ddx1-specific antibody to immunoprecipitate Ddx1-binding RNAs from Schneider2 cell lysates. Using this approach, we were able to efficiently co-immunoprecipitate Ddx1 protein, as determined by western blotting, along with Sirup mRNA, but not Ddx1 mRNA, as detected by RT-PCR (Fig. 8C-E). Ddx1 mRNA was included as a negative control to ensure that the interaction between Sirup mRNA and Ddx1 protein was specific, and not the result of non-specific RNA interaction.

Next, we tested whether there is genetic interaction between Ddx1 and Sirup. We used the UAS-GAL4 system (Actin5C-GAL4 and UAS-Sirup RNAi \( \left[ P(GD14644)\text{v36437}\right] \) to generate flies that expressed a Sirup siRNA construct in Ddx1 normal, Ddx1 heterozygous and Ddx1 null flies. Sirup knock-down in a control background reduced Sirup RNA levels by approximately 75% (Fig. 8F), and generated viable, fertile flies that displayed no observable phenotypes. Sirup knock-down in both Ddx1 heterozygous and null backgrounds resulted in lethality in the majority of progeny. Although 50% of the progeny were expected to express Sirup siRNA, we found that the majority of Ddx1 mutant progeny generated from these crosses did not contain the Act5C-GAL4 transgene (with 3 mutant progeny containing both the Act5C-GAL4 driver and the UAS-Sirup transgene versus 148 containing only the UAS-Sirup transgene) (Fig. 8G). Of note, the two Ddx1 null/Sirup knock-down progeny survived less than 1 day post eclosion, whereas the single Ddx1 heterozygous/Sirup knock-down progeny survived 14 days post eclosion.

4. Discussion

While human DDX1 has been implicated in numerous biological processes in cultured cells, we still have a poor understanding of its role in animal development. Towards this end, we generated a Ddx1 null mutant in Drosophila. In contrast to a report listing a previous Ddx1 mutation (Ddx1\(^{9W1} \)) as lethal (Zinsmaier et al., 1994), we found that Ddx1 null flies were viable. Although our results suggest that Ddx1 is not essential for embryonic development, it is important to note that significant levels of maternal Ddx1 protein are detected up to the 2nd instar stage. As Ddx1 levels are highest in early embryogenesis (Rafti et al., 1996), this maternal contribution is likely sufficient to fulfill the need for Ddx1 in early development, postponing the appearance of Ddx1 phenotypes until later stages in Ddx1\(^{AX/AX} \) flies.

The Ddx1\(^{9W1} \) mutant described by Zinsmaier et al. (Zinsmaier et al., 1994) was generated using a P-element associated with CSP located downstream of Ddx1. The authors indirectly inferred that the described allele affected Ddx1; however, unknown at the time, there is another gene between Ddx1 and CSP, CG11523. A subsequent report showed that disrupted CG11523 is associated with a lethal phenotype (CG11523\(^{Dd92S} \)) (Mummery-Widmer et al., 2009). It is, therefore, likely that the effect attributed to Ddx1 mutation in 1994 was actually due to disruption of CG11523. Alternatively, the lethal phenotype observed by Zinsmaier et al. may be the result of a truncated Ddx1 product, as the location of the P-element used to generate the Ddx1\(^{9W1} \) allele is such that deletions would likely result in a truncation. A truncated gene product
Fig. 8. **Interactions between Ddx1 and Sirup.** (A) Pictograph showing the structure of Sirup and the variable splice junction site (top – transcript expressed in control flies; bottom – transcript expressed in Ddx1<sup>AX/AX</sup> flies). (B) RT-PCR analysis of control and null larvae and adult flies. A spliced Sirup product is only observed in Ddx1 null animals. (C) IP using anti-Ddx1 antibody demonstrating that almost all Ddx1 protein is retained in the immunoprecipitate. (D, E) Left panels – RT-PCR analysis of RNA co-immunoprecipitated with Ddx1 or IgG. Right panels – RT-PCR of cDNA generated from control flies used as a positive control for the PCR reaction. Sirup mRNA results indicate that Sirup RNA, but not Ddx1 RNA, is pulled down with Ddx1 protein. (F) RT-PCR analysis shows reduced Sirup RNA levels in Sirup knock-down adult flies. (G) Progeny generated from y<sup>1</sup> w<sup>1</sup>; Actin5C-Gal4/CyO; Ddx1<sup>AX/TM3, Sb<sup>y1w<sup>1</sup>; UAS-Sirup-RNAi; Ddx1<sup>AX/TM3, Ser GFP</sup> crosses. Expected ratios of 2:1 for heterozygous to homozygous mutant Ddx1 and 1:1 for Sirup knock-down to CyO. A significant reduction in the number of Sirup knock-down flies was observed. Chi square analysis comparing observed distribution to expected distribution was used to determine significance.
could cause a divergent phenotype from that of a null allele. We are unable to test these possibilities experimentally, as the original stock is no longer available.

Although Ddx1 null flies survive to adulthood, it is clear that Ddx1 is involved in a wide spectrum of developmental functions. Ddx1 null flies are significantly reduced in size, developmentally delayed and out-competed by heterozygous animals when raised in crowded conditions. In addition, Ddx1 null flies are infertile due to aberrant gametogenesis, with both oogenesis and spermatogenesis being affected. We observed significantly smaller ovaries containing few to no mature eggs in Ddx1 null flies. One possible explanation for reduced ovary size is disruption of stem cell maintenance and/or division; however, mutants in which the stem cell pool is affected typically display reduced numbers of early egg chambers (Li et al., 2009; Shen et al., 2009). Ddx1 null ovaries appear to have normal numbers of egg chambers, suggesting that regulation of stem cell niche is not the cause of the small ovary phenotype.

We also observed autophagic egg chambers in ovaries isolated from Ddx1 null flies. Autophagic egg chambers, in addition to small body size, are indicative of metabolic stress and starvation conditions (Barth et al., 2011; Edgar, 2006; McCall, 2004). The phenotypes observed in Ddx1 null flies therefore suggest a role for Ddx1 in metabolic function and/or regulation, with absence of Ddx1 phenocopying nutrient deficiency rather than stem cell deficiency. While our attempts to generate a transgenic Ddx1 over-expression line to rescue this phenotype were not successful, we were able to confirm comparable phenotypes in flies carrying the Ddx1null allele over a deficiency that encompasses the Ddx1 locus.

In addition to autophagic egg chambers, we also noted mis-patterned mature eggs lacking dorsal appendages in Ddx1 null ovaries. Dorsal appendage formation is a complicated process involving more than 60 genes (Berg, 2005). We performed immunofluorescence analysis of Gurken localization, a key factor in dorsal appendage development, and observed loss of Gurken at the anterior dorsal corner in stage 8-9 egg chambers. Our RNA deep-sequencing did not identify any of the well-characterized Gurken localization factors as being differentially expressed in Ddx1 control versus null flies. Thus, in Ddx1 null flies, disruption of gurken translation, rather than gurken RNA/protein localization, may underlie the absence of Gurken protein in the anterior dorsal corner. Intriguingly, it has been shown that the DNA damage response and small RNA processing are required for proper Gurken translation (Chen et al., 2007; Klattenhoff et al., 2007). Human DDX1 has previously been implicated in both the DNA damage response and small RNA processing (Han et al., 2014; Li et al., 2008). The patterning defect observed in the few Ddx1 null eggs that do not undergo autophagy may therefore represent a secondary phenotype unrelated to metabolic disruption.

In keeping with a role for Ddx1 in metabolism, we found reduced p56k levels in Ddx1 null flies. p56k levels are indicative of TOR signalling, which is modified by a number of pathways including insulin signalling and nutrient sensing (Edgar, 2006). Signalling through TOR and p56k drives protein synthesis and cell growth (Zhang et al., 2000). Our RNA deep sequencing data indicate that neither TOR nor S6k mRNA levels are significantly affected in Ddx1null flies. RNA levels of major upstream regulators of the TOR pathway, including Rheb, Tsc1 and gigas (DTSC2) are also unchanged. The consistent levels of these key signalling factors suggest that the absence of Ddx1 does not directly affect the transcripts encoding proteins that participate in TOR signalling. Rather, absence of Ddx1 may modulate the function of RNA targets that function upstream of the TOR signalling pathway.

In contrast to other DEAD box proteins that have been directly implicated in Drosophila oogenesis (Cauchi, 2012; Johnstone et al., 2005; Styhler et al., 1998; Tomancak et al., 1998), we propose that the infertility observed in Ddx1 null females is due to disrupted metabolism. Altered metabolism also provides a possible explanation for the small number of progeny generated by Ddx1null/Df (3L)ED230 females, as we observed slightly higher p56k levels and slightly larger body size in these mutants compared to the completely sterile Ddx1AX/AX females. Thus, Ddx1null/Df (3L)ED230 flies may have slightly higher metabolic function than Ddx1AX/AX flies. This may be due to second-site alleles that are different between the Ddx1null and Df(3L)ED230 genetic background.

Spermatogenesis can also be affected by starvation conditions, with a noted reduction in the number of germinal stem cells in starved males (McLeod et al., 2010). In contrast to the phenotype observed in Ddx1 null ovaries, aberrant sperm individualization, the phenotype observed in Ddx1 null males, has not been associated with metabolic disruption. Following meiosis, spermiogenesis, the last stage of spermatogenesis, involves reshaping interconnected spermatids into individual mature sperm (Fabian and Brill, 2012). Spermiogenesis is characterized by gross morphological changes to the cell and mitochondria in particular. Developing spermatids in Ddx1 null males appear to undergo spermatid elongation, but fail to individualize. Instead, they fall out of the developing sperm bundle prematurely and seminal vesicles remain devoid of mature sperm.

Several DEAD box proteins have been associated with defects in sperm development. RecQ5 and Belle are both required for early spermatogenesis (Johnstone et al., 2005; Sakurai et al., 2014) and mutation of Rm62 has also been shown to cause male sterility (Buszczak et al., 2007). While the cause of sterility in Rm62 mutants has not been identified, mutation of Blanks, a binding partner for Rm62, results in spermiogenesis defects similar to those observed in Ddx1 null males (Gerbasi et al., 2011). As disrupted sperm individualization has not been linked to metabolic function, we postulate that this aspect of the Ddx1 null phenotype is the result of disruption of distinct RNA targets from those involved in metabolic function.

It has been previously established that tRNA synthesis is regulated by TOR signalling (Ciesla and Boguta, 2008) and human DDX1 has been recently identified as a tRNA splicing factor (Popow et al., 2011; Popow et al., 2014). As our analysis did not show changes in levels of spliced and unspliced tRNAs in Ddx1 null flies, it would appear that Drosophila Ddx1 is not essential for tRNA splicing. However, our northern blot analysis is only a snapshot of tRNA levels at the time of RNA isolation. Popow et al. showed that human DDX1 is necessary for efficient cycling of the Rtc8-guanylate intermediate required for tRNA splicing in vitro (Popow et al., 2014). It is possible that tRNA generation efficiency is reduced in Ddx1AX/AX flies, but a compensatory mechanism is coming into play. Alternatively, the reduction in efficiency of tRNA splicing may not be sufficient to reduce their overall steady state levels and/or our method of analysis may not be sufficiently sensitive to detect the changes in tRNA levels.

Our RNA deep sequencing analysis revealed changes in the levels of 333 RNA molecules in Ddx1AX/AX larvae. In addition, we identified a number of transcripts that were differentially spliced in Ddx1 control versus null flies. We verified that Sirup mRNA is uniquely spliced in Ddx1AX/AX larvae and up-regulated. We were also able to show that Ddx1 protein binds Sirup mRNA, which suggests that this splice site modification is due to a direct interaction with Ddx1, rather than being an indirect downstream consequence of the loss of Ddx1. Previous work has demonstrated that loss of Sirup results in shortened life span and neurodegeneration (Van Vranken et al., 2014). In agreement with our results, Sirup has also previously been shown to be up-regulated in response to starvation conditions (Erdi et al., 2012). We propose a role for Sirup in limiting metabolism during stress conditions. Sirup has recently been identified as the Drosophila homolog of...
yeast Sdh8 (Van Vranken et al., 2014). Similar to its yeast homologue, Drosophila Sirup may be required to stabilize the succinate dehydrogenase holocomplex, and enhance succinate dehydrogenase activity.

We observed spliced Sirup mRNA only in the absence of Ddx1, and propose that Ddx1 plays a role in the repression of splicing under normal conditions. This is in line with observations that human DD1X is required for efficient trafficking of unspliced viral RNA genomes (Edgcomb et al., 2012; Fang et al., 2005; Robertson-Anderson et al., 2011). There are several possible mechanisms for splicing repression by Ddx1. A simple explanation could be that Ddx1 is remodeling RNA secondary structure into a form that is refractory to splicing. Alternatively, recent studies have shown a strong relationship between antisense non-coding RNAs (ncRNAs) and alternative splicing in humans (Morrissy et al., 2011). As Ddx1 has RNA/RNA unwinding activity, Drosophila Ddx1 may act by unwinding dsRNA formed by ncRNAs bound to mRNA molecules. While it is still not clear whether ncRNAs promote or repress specific splice sites, it is possible that they may act in either fashion, depending on where they bind to target RNAs.

Although the relationship between Ddx1-dependent splice regulation of Sirup mRNA and the metabolic disruption observed in Ddx1 null flies remains a matter of speculation, Sirup’s known role in mitochondrial function provides a possible explanation for the epistatic lethality observed in Ddx1 null/Sirup knock-down flies. We propose that Sirup’s unspliced form is required for steady state mitochondrial function. The unspliced form of Sirup retains an early stop codon that produces a transcript that encodes a short 36 aa product missing conserved domains required for succinate dehydrogenase activity. However, there is evidence suggesting that this stop codon undergoes read-through by ribosomes, thereby generating a 145 aa length protein in spite of the stop codon (Dunn et al., 2013). Under stress conditions, suppression of Ddx1 activity may promote splicing of Sirup mRNA thereby generating a shorter 118 aa product that has reduced succinate dehydrogenase activity. The resulting slowdown in mitochondrial function would reduce resource usage by the cell, allowing it to survive periods of nutrient scarcity. Under this model, the ob- function would reduce resource usage by the cell, allowing it to survive periods of nutrient scarcity. Under this model, the ob- function would reduce resource usage by the cell, allowing it to survive periods of nutrient scarcity. Under this model, the ob- function would reduce resource usage by the cell, allowing it to survive periods of nutrient scarcity. Under this model, the ob- function would reduce resource usage by the cell, allowing it to survive periods of nutrient scarcity. Under this model, the ob- function would reduce resource usage by the cell, allowing it to survive periods of nutrient scarcity. Under this model, the ob- function would reduce resource usage by the cell, allowing it to survive periods of nutrient scarcity. Under this model, the ob- function would reduce resource usage by the cell, allowing it to survive periods of nutrient scarcity. Under this model, the ob- function would reduce resource usage by the cell, allowing it to survive periods of nutrient scarcity. Under this model, the ob- function would reduce resource usage by the cell, allowing it to survive periods of nutrient scarcity. Under this model, the ob- function would reduce resource usage by the cell, allowing it to survive periods of nutrient scarcity. Under this model, the ob- function would reduce resource usage by the cell, allowing it to survive periods of nutrient scarcity. Under this model, the ob- function would reduce resource usage by the cell, allowing it to survive periods of nutrient scarcity. Under this model, the ob- function would reduce resource usage by the cell, allowing it to...


