dMARCH8, a Drosophila ubiquitin E3 ligase, regulates polymodifications of tubulin

- 2 in the spermiogenic pathway
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11 ABSTRACT

Ciliary stability and function are regulated by the covalent addition of polyglutamate and polyglycine chains to axonemal tubulin subunits. The *Drosophila* gene CG13442 encodes a predicted ubiquitin E3 ligase involved in the regulation of tubulin glycylation and glutamylation. Homologous to mammalian MARCH8, CG13442/dMARCH8 is required for male fertility. Sperm in *dMARCH8* mutant testes appear to undergo a normal individualization process but fail to be transferred to the seminal vesicle. This phenotype is very similar to that of mutants in the *Ntl* gene, shown here to be a glycine transporter using a [³H]glycine uptake assay. Mutations in *dMARCH8* are associated with a reduction of both polyglutamylation and polyglycylation of sperm tubulin. Polyglutamylation of tubulin is significantly increased in the *Ntl* background, and recovers to wild-type levels in the *Ntl* double mutant background, indicating that glycine and glutamate compete for some common site(s) on tubulin molecules in this system. By analogy to the regulation of the mammalian glycine transporter GlyT2 through ubiquitin-mediated trafficking between the plasma membrane and endosome, *dMARCH8* may target *Ntl* and glutamate transporters, or other upstream regulators of these proteins.

Introduction

Amino acids and their chemical derivatives serve as signaling molecules in several biological contexts. Glycine and glutamate function as major inhibitory and excitatory neurotransmitters, respectively, in the central nervous system (Kandel et al., 2013). At the subcellular level, these same two amino acids regulate ciliary function and stability through a mechanism involving their competition for covalent addition of polyglycine and polyglutamate chains to tubulin subunits incorporated into stable microtubule-based structures (Raunser and Gatsogiannis, 2015). The activity of glycine transporters, controlling the levels of glycine both inside and outside of the cell, can therefore result in behavioral effects and ciliary dysfunction (Boison, 2016)

Here we use the *Drosophila* spermiogenesis system to examine the regulation of tubulin polymodifications. *Drosophila* spermiogenesis provides an opportunity for genetic analysis of many different cellular subsystems (Fabian and Brill, 2012). These include the differentiation of mitochondria (Hales and Fuller, 1997;Politi et al., 2014), polarization of the spermiogenic cyst (Wei et al., 2008), the radical restructuring of spermatid nuclei (Kost et al., 2015), and the complex process of sperm individualization, in which individual sperm are finally resolved from the syncytium in which they have developed (Arama et al., 2007;Arama et al., 2003;Fabrizio et al., 1998;Tokuyasu et al., 1977) This complexity is reflected in the large number of genes that are mutable to a male-sterile phenotype, and indicates that genes contributing to any of the many different cellular subsystems---including the polymodifications of tubulin---may be accessible through the analysis of male-sterile mutations (Wakimoto et al, 2007).

Because there is very little transcription in spermiogenic cysts(Barreau et al., 2008) posttranscriptional and posttranslational regulatory mechanisms are especially important in this process (Karr, 2007). Ubiquitination is one of these processes, affecting many different aspects of spermiogenesis (Richburg et al., 2014). Although the ubiquitin system is known primarily for marking proteins for degradation (Ciechanover, 2005;Zhi et al., 2013), recent studies indicate that ubiquitination can affect many other aspects of protein function, including intracellular trafficking (de Juan-Sanz et al., 2011),

modulation of protein-protein interactions (Yang et al., 2010), and modulation of transcription, DNA repair and transmembrane signaling (Metzger et al., 2012).

The substrate-specificity of a ubiquitination system is generally controlled by E3 ubiquitin ligases. Here we identify an E3 ubiquitin ligase, *dMARCH8*, homologous to mammalian MARCH8. *dMARCH8/CG13442* function is required for the full complement of polyglycylation and polyglutamylation of sperm tubulin in the *Drosophila* testis. We show that the *Drosophila Neurotransmitter transporter-like (Ntl)* gene, previously found to exhibit a similar phenotype to *dMARCH8* mutants (Chatterjee et al., 2011), encodes a glycine transporter using a [³H]glycine uptake assay. Loss of *Ntl* function results in a large increase in polyglutamylation levels, indicating that *Ntl* and *dMARCH8* activities contribute to the balance of glutamylation and glycylation of sperm tubulin.

Materials and Methods

Fly Husbandry

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- 74 Flies were raised on standard cornmeal molasses agar at 25°C. Unless otherwise
- 75 mentioned, all stocks were from the Bloomington Stock Center. Males were tested for
- 76 fertility by mating in groups of 4–5 with an equal number of virgin females. Generally,
- 77 w^+ or y^+ males were mated with yw females, with the recovery of w^+ or y^+ daughters in
- 78 the F1 generation confirming fertility. Genetic constructions were carried out using
- 79 standard *Drosophila* genetics as in Greenspan (Greenspan RJ, 1997). All experiments
- were carried out in the yw genetic background.

Generation of dMARCH8 mutants

- 82 The dMARCH8 transcript/CDR is in the 57B region on the 2R arm of the Drosophila
- chromosome. The Mi{MIC}(Minos Mediated Integrated Cassette) $dpr^{Ml0657l}$ transposon
- 84 carrying the y^+ marker was generated by the *Drosophila* Gene disruption project
- 85 (GDP)(Venken et al., 2011)The transposon was mobilized by crossing the flies carrying
- 86 the MiMIC insertion to flies carrying stable heat shock transposase $(y^1 w^*; sna^{Sco} / SM6a,$
- 87 $P\{w^{+mC}=hsILMiT\}$ 2.4 (Metaxakis et al., 2005; Venken et al., 2011). Three broods from
- each cross were generated by transferring flies to new bottles on the 3rd and 5th days after
- 89 the cross was initially set up. Bottles were subjected to heat shock (39°C, 90 minutes) on
- 90 the third, 5^{th} and seventh day of each brood. Chromosomes that lost the y^+ markers
- 91 carried by the transposon were recovered and established in balanced stocks using
- 92 standard *Drosophila* genetics, then screened for new male-sterile mutations produced by
- 93 imprecise excision of the Mi{MIC} transposon.

RNA isolation and qRT-PCR

- 95 Total RNA was isolated using TRI reagent (Sigma) according to the manufacturer's
- 96 recommendations. RNA was extracted from males, females, testes, and heads and
- 97 concentration was determined by measuring its absorbance.
- 98 One-step gRT-PCR was performed using the iTag Universal SYBR Green One-Step Kit
- 99 (BIORAD) according to the manufacturer's recommendations. The BIORAD MyiO
- Real-Time PCR Detection System was programmed as follows: 50°C for 10 min, 95°C

- 101 for 5 min followed by the amplification steps of 95°C for 15 sec, 56°C for 30 seconds,
- 102 72°C for 30 sec. 45 cycles of PCR were run for all samples followed by 72°C for 15 min,
- 103 57°C for 10 min and held at 4°C overnight. dMARCH8 qRT primers amplified a 112 bp
- fragment, while rp49 control primers amplified a 154 bp fragment. Primers for this and
- all subsequent PCR-based molecular biology are specified in Table 1.

Deletion PCR

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- 107 Genomic DNA was isolated from males according to the Berkeley Drosophila Genome
- Project protocol (http://www.fruitfly.org/about/methods/inverse.pcr.html). 13 pairs of
- gene-specific primers were used to amplify the entire coding sequence of dMARCH8.
- 110 The thermocycler (MJ Research PTC-200 Peltier thermal cycler) was programmed as
- follows: 94°C for 2 min followed by 30 cycles of: 94°C for 1 min, 56.3°C for 1 min,
- 72°C for 1 min. After a final 5 min at 72°C, samples were held at 4°C until gel analysis.

Phalloidin assay for Individualization Complex

- 114 Testes from 0-1 d old males were crudely dissected in *Drosophila* Ringers and
- transferred immediately to a tube of Ringers on ice. Testes were then fixed for 15 min in
- 116 4% paraformaldehyde in buffer B (16.7mM KH₂PO₄/K₂HPO₄ pH 6.8, 75mM KCl,
- 117 25mM NaCl, 3.3mM MgCl₂). Following fixation, testes were rinsed three times in PTx
- 118 (PBS+0.1% Triton X-100), washed for 30 min in PTx and blocked for at least 1 h in
- blocking solution (0.01% NaAzide and 3% BSA in PTx). Staining with rhodamine-
- 120 conjugated phalloidin was for 30 min in blocking solution (3 μg/ml). Testes were then
- rinsed and washed with PTx. Testes were then finely dissected from remaining carcasses
- in 50% glycerol and then mounted in 90% glycerol. Slides were stored overnight at 4°C
- before imaging. Fluorescence images were captured by confocal microscopy (Leica TCS-
- 124 2, Exton, PA)(Fabrizio et al., 2012)

Phase squashes

- 126 Testes were dissected from 0-1-day-old young male *Drosophila* mutants (unless
- mentioned otherwise) in 1x PBS buffer and gently squashed with a coverslip before
- taking phase images with Leica DM 4500B. For phase microscopy of seminal vesicles,

wild type and mutant freshly eclosed Drosophila males were withheld from females for

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three to four days before dissecting their testes. **Protein Electrophoresis and Immunoblot** Samples were prepared from male fly testes with seminal vesicle for each genotype, from males that were withheld from females for 4–6 days in order to maximize sperm yield. Six fly testes (12 testes) worth of protein were loaded in each lane. Testes were dissected in 1× PBS, ground in 2× Laemmeli buffer (Laemmli UK., 1970) vortexed and boiled for 5 min. Samples were then spun at 15,800 g for 5 min, and supernatants were separated in a 12% SDS-Polyacrylamide gel and transferred to a PVDF membrane (Amersham, GE Healthcare, United Kingdom) using a Trans-Blot Semi-Dry transfer apparatus (BioRad, U.S.A). Membranes were incubated with primary antibodies directed against polyglycylated tubulin (Poly-G) (1:10,00) (MABS276, cloneAXO49, Millipore), polyglutamylated tubulin (Poly-E) (1:1000) (mAbGT355, Adipogen Life sciences), ubiquitin {MAB1510 clone Ubi-1 (aka 042691GS), Millipore}, TSSK2 (ab172434, Abcam) and α-tubulin (1:500), (DM1A, Sigma). Protein bands were visualized with HRP-labelled anti rabbit or anti-mouse (1:5,000) secondary antibodies followed by detection with ECL immunoblot detection kit (Pierce, U.S.A). Loading control was αtubulin. All statistical analyses were made using two-tailed Students t-test in Microsoft Excel. The average values of relative intensity (Target antibody intensity/anti-tubulin) were plotted for each genotype. Intensities were calculated using Image J software (Rasband, W.S., ImageJ, U. S. National Institutes of Health, Bethesda, Maryland, USA, http://rsb.info. nih.gov/ij/, 1997–2009). [3H]glycine uptake assays Human GlyT1, GlyT2 and *Drosophila* Ntl cDNAs were cloned into pRK5myc or pTMR vectors as previously described (Carta et al., 2012). HEK293 cells were grown in minimal essential medium (Earle's salts) supplemented with 10% (v/v) FCS, 2 mm Lglutamine, and 20 units/ml penicillin/streptomycin in 5% CO₂, 95% air. The cells were plated on poly-D-lysine-coated 24-wells plates (Nunc), grown to 50% confluence, and transfected with 1 µg of total pRK5myc-hGlyT1, pRK5myc-hGlyT2, pRK5myc-Ntl or

159 pTMR-Ntl plasmid DNAs using Lipofectamine LTX reagent (Invitrogen). After 24 h, the 160 cells were washed twice with prewarmed buffer (118 mM NaCl, 1 mM NH₂PO₄, 26 mM 161 NaHCO₃, 1.5 mM MgSO₄, 5 mM KCl, 1.3 mM CaCl₂, 20 mM glucose) pre-equilibrated with 5% CO₂, 95% air. After 2 min, the cells were incubated for 5 min in 0.1 μCi/ml 162 [³H]glycine (60 Ci/mmol; PerkinElmer Life Sciences) at a final concentration of 300 μM. 163 The cells were rinsed twice with ice-cold buffer pre-equilibrated with 5% CO₂, 95% air 164 and then digested in 0.1 M NaOH for 2 h. The samples were used for scintillation 165 166 counting and for determination of protein concentration using the Bradford reagent (Bio-Rad). [3H]glycine uptake was calculated as nmol/min/mg of protein and expressed as 167 percentages of that in control cells transfected with the empty expression vector. All 168 169 statistical comparisons used an unpaired Student's t-test.

Generation of Ntl-dMARCH8 double mutant

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Ntl-dMARCH8 double mutants were generated by crossing Ntl/CyO males to dMARCH8^{7A}/CyO females. Straight-winged female progeny obtained were crossed to yw, Gla/Sm6a males. Isomales from this cross were crossed again to establish balanced stocks. Males from these established stocks were checked for fertility by crossing to yw female virgins. Males from sterile stocks obtained were screened for complementation with Ntl and dMARCH8 mutants.

Results

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- Amino acid sequence alignment of dMARCH8 (CG13442) with human MARCH8-
- 180 E3 ubiquitin Ligases belonging to the RING protein family
- 181 A ClustalW alignment of the predicted dMARCH8 protein sequence with human
- 182 MARCH8 is shown in Figure 1A. RING domain considered critical for E3 ubiquitin
- ligase function (Samji et al., 2014) are conserved in the predicted dMARCH8 protein.
- 184 The TMpred algorithm (Hofmann and Stoffel, 1993) predicts two transmembrane
- domains for dMARCH8, characteristic of the MARCH family (Samji et al., 2014), whose
- 186 RING domains are most closely related to the Ringv and PHD domains (Deshaies and
- 187 Joazeiro, 2009).

Generation of *dMARCH8* mutants

- The dMARCH8 gene lies within an intron of the dpr gene, where it is expressed from the
- 190 opposite strand. dMARCH8 mutants were generated by mobilizing the
- 191 Mi{MIC} dpr^{M106571} near the 5' end of the dMARCH8 gene, crossing it into a genetic
- background expressing the minos transposase (Venken et al., 2011) under heat shock
- 193 control (Metaxakis et al., 2005). By standard fly genetics, chromosomes which had lost
- the $vellow^+(y^+)$ markers associated with the Mi{MIC} transposon were recovered and
- established in balanced stocks. Transposon excisions are often imprecise and deletions of
- varying size, usually extending from the insertion site in either direction, are often
- recovered at significant frequencies (Zhang and Spradling, 1993). A schematic of the
- 198 dMARCH8 locus showing the starting insertion, the mating scheme used to mobilize the
- transposon and identify new male-sterile mutations, and the location of primer pairs used
- 200 to assay for deletions in the resulting male-sterile stocks is shown in Figure 1B and 1C.
- Of 64 chromosomes that were observed to have lost the y^+ marker, 5 were found to carry
- 202 new male sterile mutations defining a single complementation group.

Deletion PCR analysis of CG13442/dMARCH8 mutants

- The mutant alleles were screened with nested gene-specific primers spanning the site of
- 205 the transposon insertion (Materials and Methods and Fig. 1B). Genomic DNA from
- 206 homozygous males carrying each of the 5 male-sterile alleles was used to probe for

- 207 changes in the chromosome structure by PCR. For all of the male-sterile mutants, PCR
- 208 amplicons including portions of the gene and flanking regions on either side were
- 209 missing or produced products with altered size as compared to wild-type DNA (Table 2).
- These results indicated that the deletion carried by the dMARCH8^{7A} mutant removes the
- 211 sequences encoding the RING domain and both transmembrane domains (Table 2).
- Furthermore, qRT-PCR analysis of this mutant revealed a significant reduction (80 +/-
- 213 5% in dMARCH8 expression (Fig. 2A). The dMARCH8^{7A} allele was used for the
- 214 remaining phenotypic analyses.

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Testis Specificity of dMARCH8 expression

- 216 qRT-PCR analysis confirmed that dMARCH8 expression was male-specific and limited
- 217 to the testes (Fig. 2B and C). Expression in females was 90 +/- 5 % lower than in males,
- and in the adult heads and carcass of males, it was 98 +/- 1% lower than in the testis.

Rescue by Germline Transformation

- We used a P-element based construct for germline transformation with CG13442/
- 221 dMARCH8, to confirm that the male sterile phenotype in our mutants is caused solely due
- 222 to disruption of dMARCH8. For this we used a pTMR-dMARCH8 construct, which
- contains a full-length cDNA for CG13442 (DGRC clone AT03090) cloned downstream
- of the β2T-tubulin transcriptional control sequences in the *Drosophila* transformation
- vector pTMR (Clark et al., 2006; Huh et al., 2004). This construct provides strong germ
- cell-specific transcription in developing sperm (Kaltschmidt et al., 1991; Kemphues et al.,
- 227 1982). A BAC genomic construct CH322-140N02 (BACPAC resources,
- bacpac.chori.org) extending from 9.8 kb upstream of the dMARCH8 transcript to 8.5 kb
- beyond the 3' end also rescued the dMARCH8 mutant phenotype. Two independent
- insertions of this construct were tested and they both rescued the mutant. All the males
- 231 homozygous for the dMARCH8^{7A} deletion and carrying pTMR-dMARCH8 or BAC-
- 232 *dMARCH8* constructs were fertile. Results from the rescued line are presented in Fig.
- 233 2D. The male-sterile mutations obtained after mobilization of the Mi{MIC} transposon
- 234 therefore result from disruption of *dMARCH8*.

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Spermiogenic-defective phenotype of *dMARCH8* mutants dMARCH8^{7A} / dMARCH8^{7A} males produced elongated spermiogenic cysts (Fig. 3), but we did not detect mature sperm in the seminal vesicle (SV) (Fig. 3B, arrows). In contrast, wild-type control seminal vesicles were filled with mature sperm (Fig. 3A, arrows). No motile sperm are seen in dMARCH8^{7A} / dMARCH8^{7A} squash preparations, unlike in the wild-type controls, where dense masses of mature motile sperm were evident (Fig. 3C, arrow). Instead, an extensive mass of sperm bundles accumulated at the base of dMARCH8^{7A} mutant testes (Fig. 3B, asterisks). Presumably because of the great physical complexity of the sperm individualization process (Bazinet and Rollins, 2003; Noguchi et al., 2008; Tokuyasu et al., 1972), a preponderance of male-sterile mutations produce elongated cysts that fail to mature into individual sperm (Wakimoto et al, 2007). In most of these cases, the individualization complex either fails to form, does not progress, or breaks down during transfer along the length of the cyst (Fabrizio et al., 1998). When dMARCH8^{7A} mutant Drosophila testes were stained with rhodamine-conjugated phalloidin, we observed normal development and movement of the actin cones of the individualization complex (Fig. 3F). Waste bag deposition in the distal end of the testis also appeared normal, further indicating that the individualization complex was successfully navigating the entire length of the cysts (Fig 3F). These observations are similar to the phenotype seen in Neurotransmitter transporter-like (Ntl) mutants, a putative glycine transporter homologous to human glycine transporters GlyT1 and GlyT2 (Chatterjee et al., 2011). Demonstration of Glycine transport activity from Ntl expression in mammalian cells To test whether the *Drosophila* Ntl protein was capable of glycine uptake, we tested the capacity of recombinantly-expressed Ntl to mediate the uptake of [3H]glycine, using human GlyT1 and GlyT2 cDNAs (Carta et al., 2012) as controls. As expected, both pRK5-GlyT1 and pRK5-GlyT2 constructs resulted in significant increases in [3H]glycine

uptake compared to vector only controls (pRK5-GlyT1: 345 ± 42; pRK5-GlyT2 299 ±

38). Both Drosophila Ntl expression constructs also resulted in statistically significant

264 [3 H]glycine uptake (pRK5-NTL: 245 ± 24; pTMR-NTL: 344 ± 59) confirming that Ntl is 265 a fully-functional glycine transporter (Fig. 4). 266 Reduction of ubiquitination of testis proteins in *dMARCH8* mutants 267 Since dMARCH8 is a member of the E3 ubiquitin ligase family, we assayed dMARCH8 268 mutant males for total ubiquitination of testis proteins using an anti-ubiquitin antibody 269 (Kane et al., 2014). The results from three independent experiments, using α -tubulin as a 270 loading control, showed an apparent reduction in ubiquitination signal in mutant samples 271 relative to wild-type controls (Fig. 5A). Two distinct ubiquitinated bands of ~50 and 77 kDa present in the testes of dMARCH8^{7A}/+ flies are missing in the testes of homozygous 272 dMARCH8^{7A} flies. A slight reduction in several other bands (~65 and 100 kDa) was also 273 consistently observed in the *dMARCH8*^{7A} mutant. 274 275 Reduction of tubulin polyglycylation and polyglutamylation in dMARCH8 mutants 276 The tubulin in *Drosophila* sperm axonemes undergoes a variety of posttranslational 277 modifications (Janke, 2014; Kierszenbaum, 2002;Wloga and Gaertig, 278 2010)Polyglycylation of tubulins is required for the stability of ciliary and flagellar 279 axonemes and other long-lived tubulin-based structures (Bre et al., 1996;Bressac et al., 280 1995;Rogowski et al., 2009a). Polyglutamylation is known to regulate beating behavior 281 in motile cilia via the regulation of flagellar dynein motors (Ikegami et al., 2010; Janke et 282 al., 2005; Kubo et al., 2010; Pathak et al., 2007; Suryavanshi et al., 2010). In addition, 283 polyglycylation and polyglutamylation have been shown to affect male fertility 284 (Chatterjee et al., 2011; Lee et al., 2013). 285 286 The spermiogenic defect of dMARCH8 mutants at the microscopic level is similar to that 287 seen in Ntl, which encodes a glycine transporter in whose absence polyglycylation of 288 testes tubulin is significantly reduced and sperm fail to be transferred to the seminal 289 vesicle. We therefore analyzed the levels of polyglycylated and polyglutamylated tubulin 290 in testis protein samples using antibodies directed against poly-G and poly-E. 291 Quantitation by scanning the results from six independent experiments, using α -tubulin as 292 a loading control showed an average of 50% reduction in poly-G signal in the dMARCH8^{7A} mutant samples relative to wild-type controls (Fig 5B). Similarly, for the 293

poly-E signal, quantitation from three independent experiments exhibited an average of

70% reduction in *dMARCH8*^{7A} mutants compared to wild-type controls (Fig. 5C).

Polyglycylation and polyglutamylation in a Ntl-dMARCH8 double mutant

Since the *Ntl* mutant exhibits a similar phenotype to the *dMARCH8* mutant, and it also shows a predicted interaction (STITCH database) with dMARCH8, we analyzed the *Ntl*-

dMARCH8 double mutant for polyglycylation and polyglutamylation levels of tubulin in

three independent experiments. We observed that in both the Ntl mutant and the Ntl-

dMARCH8 double mutant, poly-G levels decrease while the poly-E levels increase (Fig.

5D,E and F). Perhaps most strikingly, loss of Ntl in the dMARCH8 genetic background

restores levels of polyglutamylation to wild-type levels (compare Fig. 5C and 5F). In this

system, polyglutamylation appears to be very sensitive to glycine levels, consistent with

observations by others that glutamate and glycine compete with each other for common

site(s) on tubulin (Bulinski, 2009;Rogowski et al., 2009b;Wloga et al., 2009).

Discussion

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308 Functional spermatogenesis is particularly dependent on ubiquitin-regulated protein

function and stability (Mukhopadhyay and Riezman, 2007; Richburg et al., 2014). This is

evident from the spermatogenesis-specific functions of a number of E3 ligases: Bruce and

311 Cullin3 in *Drosophila* have been studied for their role in sperm individualization (Arama

312 et al., 2007; Arama et al., 2003; Kaplan et al., 2010; Wang et al., 2006). RNF8 and E3^{histone},

found in mice and rats respectively, are reported to be involved in the histone degradation

that occurs during histone to protamine transition in spermatid nuclei in rats (Liu et al.,

315 2005;Lu et al., 2010). The E3 ligases Cul4A (Kopanja et al., 2011;Yin et al., 2011), Itch

(Dwyer and Richburg, 2012) and Siah1a (Dickins et al., 2002) play roles during germ cell

meiosis and Cullin3^{testis} (Arama et al., 2007; Kaplan et al., 2010) and MEX (Nishito et al.,

2006) are required during germ cell apoptosis. MARCH7 (Membrane-Associated Ring-

<u>CH</u>) was found to be involved in spermiogenesis by regulating the structural and

functional integrity of the head and tail of developing spermatids (Zhao et al., 2013).

321 MARCH10 was found to be essential for spermatid maturation (Iyengar PV, Hirota T,

Hirose S, Nakamura N., 2011). MARCH11 plays a role in ubiquitin-mediated protein

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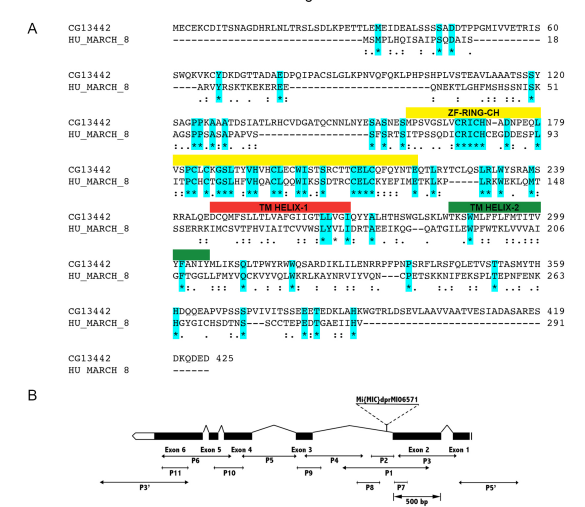
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sorting in TGN-MVB (Trans Golgi Network-multivesicular bodies) transport in developing spermatids (Morokuma et al., 2007; Yogo et al., 2012). Indeed, of all organs examined in rats, ubiquitination has been reported to be highest in testis (Rajapurohitam et al., 2002). Combined with the observed genetic sensitivity of the spermiogenic process, these considerations indicate that details of cellular ubiquitin function may be accessible through the characterization of male-sterile mutations in E3 ligase genes. Studies of mammalian spermatogenesis have implicated ubiquitination in the regulation of multiple spermiogenic stages, including nuclear condensation, acrosome formation and membrane transport (Nakamura, 2013). Here we have observed that loss of the E3 ubiquitin ligase dMARCH8 results in a malesterile phenotype very similar to that observed in mutants of the transporter Ntl. We have also shown Ntl mediates the uptake of [3H]glycine in a cellular assay with an efficiency comparable to the mammalian glycine transporters GlyT1 and GlyT2. At least one other uncharacterized *Drosophila* SLC6 class transporter encoded by CG5549, gives a somewhat better BLAST score against human GlyT1 (42.34% identity) and GlyT2 (35.01 % identity) than Ntl (Ntl vs GlyT1 39.15%; Ntl vs GlyT2 37.6% identity) suggesting that the fly genome may encode an additional glycine transporter. Expression of CG5549 is also male biased {Flybase}, although not as strongly as that of *Ntl*. Modulation of the activity of the mammalian glycine transporter GlyT in mammalian cells is thought to be achieved via ubiquitin-dependent endocytosis of GlyT to recycling endosomes, from which it can be returned to the plasma membrane when needed (Barrera et al., 2015; de Juan-Sanz et al., 2011; de Juan-Sanz et al., 2013; Fernandez-Sanchez et al., 2009). Ubiquitylation of the GlyT is associated with internalization of the transporter, where it can where it can no longer contribute to cellular glycine uptake. In that case, one might expect constitutively higher levels of glycine in cells that had lost the ability to ubiquitinate the transporter. If the Ntl transporter was similarly regulated through ubiquitylation by dMARCH8, one might expect that loss of dMARCH8 would not result in reduced availability of glycine as a result of the intracellular sequestration of the Ntl transporter. Instead, the opposite is observed, a reduction in polyglycylation and glutamylation. This suggests that Ntl and potentially a glutamate transporter are not

themselves substrates of *dMARCH8*, but that the reduced glycylation and glutamylation results from the activity of *dMARCH8* on factors upstream of glycine/glutamate transport, or in some parallel pathway. Future studies will seek to clarify whether Ntl is a substrate of dMARCH8 and to identify and characterize any additional components of the system controlling spermiogenic tubulin glycylation and glutamylation.

Figure 1



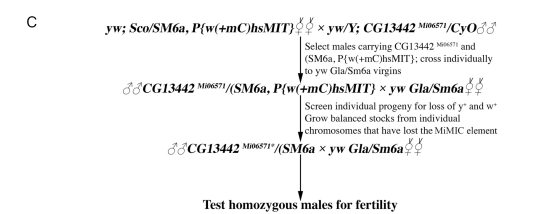


Figure 1: Amino acid sequence alignment of *Drosophila melanogaster CG13442* (dMARCH8) with human MARCH 8 using ClustalW alignment and generation of mutants in the dMARCH8 (CG13442) gene.

A) Strictly conserved residues are highlighted in blue. Conserved RING-CH domain is highlighted in yellow and transmembrane domains 1 and 2 are highlighted in red and green respectively.

B) *dMARCH8* (CG13442) transcript/CDR at 57B1 on the right arm of chromosome 2. MiMIC transposon element is inserted between Exon2 and Exon3, closer to Exon2.

C) Scheme for generation of *dMARCH8* (CG13442) deletion mutations by imprecise excision of the MiMIC transposon

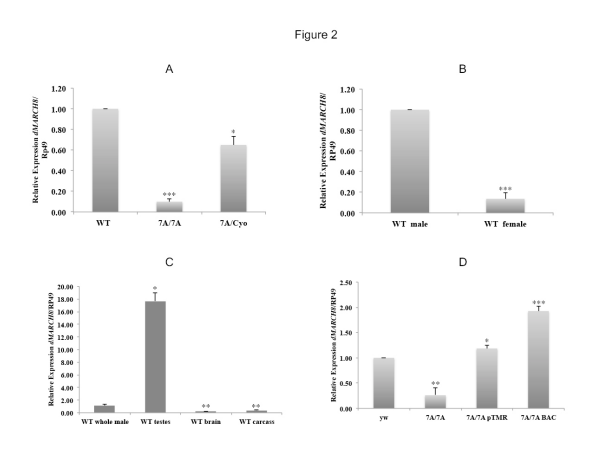


Figure 2: Analysis of dMARCH8 mRNA Expression

- 371 A) qRT-PCR analysis shows decrease in dMARCH8 expression levels in
- 372 dMARCH8^{7A}/dMARCH8^{7A} (7A/7A) mutants as compared to wild-type(yw) and
- heterozygous dMARCH8^{7A}/CyO flies. Bar, standard error of the mean. N=3 (* p<0.05, **
- 374 p<0.01, **p<0.001 by two-tailed t-test).
- B) qRT-PCR analysis compares the relative expression of dMARCH8 in wild-type (yw)
- male and female (whole adult flies). Bar, standard error of the mean. N=3 (* p<0.05, **
- 377 p<0.01, **p<0.001 by two-tailed t-test).
- 378 C) qRT-PCR products from wild-type (yw) whole male, (yw) male testes, (yw) male
- brain, and (yw) male carcass. Bar, standard error of the mean. N=3 (* p<0.05, ** p<0.01,
- **p<0.001 by two-tailed t-test).
- 381 D) dMARCH8 qRT-PCR products from dMARCH8^{7A}/dMARCH8^{7A} (7A/7A) mutant
- males carrying a pTMR-dMARCH8 cDNA construct (yw; dMARCH8^{7A}/dMARCH8^{7A};
- 383 pTMR- dMARCH8) (7A/7A;pTMR) and a genomic BAC construct (yw;
- 384 dMARCH8^{7A}/dMARCH8^{7A}; BAC-dMARCH8) (7A/7A;BAC) compared to mutant
- 385 *dMARCH8*^{7A}/*dMARCH8*^{7A} and wild-type controls (*yw*). Bar, standard error of the mean.
- 386 N=3 (* p<0.05, ** p<0.01, **p<0.001 by two-tailed t-test).

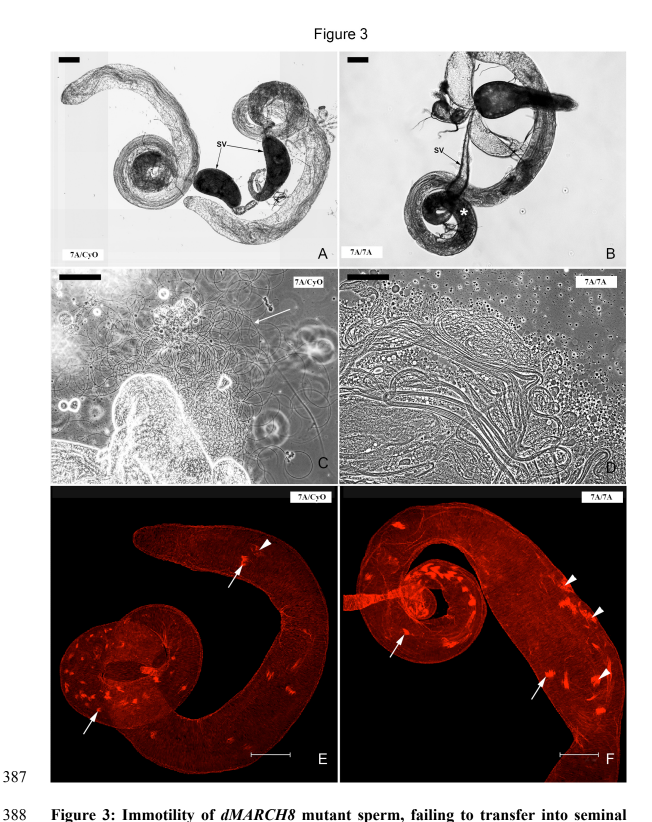


Figure 3: Immotility of *dMARCH8* mutant sperm, failing to transfer into seminal vesicles and individualization in *dMARCH8* mutants.

Panels A, B, C and D: Phase contrast images of testes from *dMARCH8*⁺ (A, C) and *dMARCH8*⁻ (B, D) males. The major phenotypic feature of the mutants is the accumulation of coiled cysts at the base of the testis (asterisks), and the empty/shrunken state of the seminal vesicle (SV) (arrows). In panel C, the letter M denotes dense masses of mature motile sperm, which is not seen in the mutants. Left hand panels: heterozygous (*dMARCH8*^{7A}/*Cyo*); right hand panels: (*dMARCH8*^{7A}/*dMARCH8*^{7A}) mutants. Bars, 100 μm.

E: Heterozygous *dMARCH8*^{7A}/*Cyo* testis stained with rhodamine-conjugated phalloidin to visualize the actin cone-based individualization complexes. Arrows mark the actin cones of the complex. Arrowheads mark the waste-bags.

F: *dMARCH8*^{7A}/ *dMARCH8*^{7A} mutant testis preparations stained with rhodamine-conjugated phalloidin to visualize the actin cone-based individualization complexes. Formation and movement of actin cones/individualization complex along the mutant cysts appears normal. Bars, 100 um.

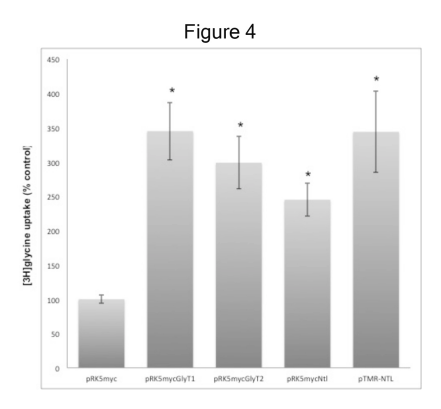


Figure 4: Drosophila Ntl is a functional glycine transporter.

Glycine uptake in HEK293 cells transiently expressing hGlyT1, hGlyT2 and *Drosophila* Ntl expressed from two different expression vectors after 5 min of incubation with $[^3H]$ glycine at a final concentration of 300 μ M. Because low levels of glycine uptake are found in HEK293 cells (Carta et al., 2012), $[^3H]$ glycine uptake was calculated as nmol/min/mg of protein and then expressed as a percentage of the empty expression vector (pRK5-myc) transfected control. The data are the means \pm S.E. (n = 7-8). Statistical comparisons were made using an unpaired Students t-test. The asterisk indicates significantly different from empty vector control (p < 0.01).

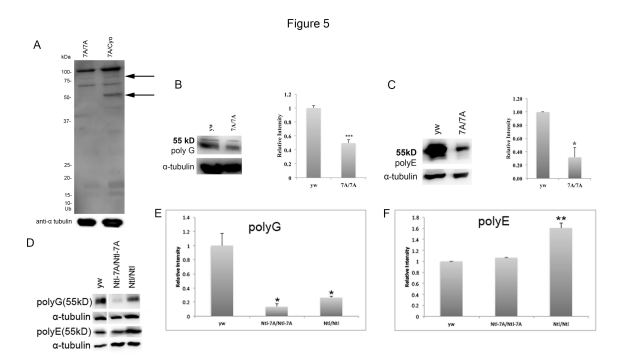


Figure 5: Western Blot Analysis of dMARCH8^{7A}/dMARCH8^{7A}, Ntl/Ntl and Ntl-dMARCH8^{7A} mutant testes.

A) Total ubiquitination levels of testis proteins are reduced in dMARCH8 mutant testes. Western blot of dMARCH87A/dMARCH87A (dMARCH8-) males compared to dMARCH87A/CyO (dMARCH8+) heterozygous males probed with anti-ubiquitin antibody.

B) Polyglycylation of tubulin is partially decreased in mutant testes. Western blot of dMARCH87A/dMARCH87A (dMARCH8-) males compared to yw (dMARCH8+/+)

probed with anti Poly-G antiserum. Quantitation of three independent replicates of the Western analysis. Bar. standard error of the mean. N=3 (* p<0.05, ** p<0.01, **p<0.001 by two-tailed t-test). C) Polyglutamylation of tubulin is partially decreased in mutant testes. Western blot of dMARCH87A/dMARCH87A (dMARCH8-) males compared to vw (dMARCH8+/+) controls probed with anti Poly-E antiserum. Quantitation of three independent replicates of the Western analysis. Bar, standard error of the mean. N=3 (* p<0.05, ** p<0.01, **p<0.001 by two-tailed t-test). (D, E, F) Polyglycylation (polyG) and polyglutamylation (polyE) levels of tubulin in Ntl/Ntl and Ntl-dMARCH87A mutant testes. Western blot showing tubulin polyglycylation and polyglutamylation levels in Ntl/Ntl (lane 3) and Ntl-dMARCH87A/ Ntl-dMARCH87A (lane 2) mutant testes compared to vw (WT) (lane 1) probed with anti Poly-G and Poly-E antiserum. Quantitation of three independent replicates of the polyG and polyE Western analysis respectively. Bar, standard error of the mean. N=3 (* p<0.05, ** p<0.01, **p<0.001 by two-tailed t-test).

Primer	Sense	Antisense
pair		
Interval	GGGCATGATCGTTGTGGAGA	AAATAAATGCATTCGCTCGC
P1		
Interval	CCGATAATCCCGAACAGTGA	GAAAAACCCCACTGGTGATG
P2		
Interval	CGTCGAACCGATCAAAGAAT	ATGCGGACTCGTAATTCAGG
P3		
Interval	TCCCCGCATTTGAATATTGT	GCGAGGTGCTAATCCAACAC
P4		
Interval	AAGGTGAGCAGCGAGAACAT	CGGAACAGACTCTGCGGTA
P5		
Interval	CCTGCTGGTAGGCATCCAGTACTA	GAAAGCCCAGCAAAATCCTT
P6		
Interval	CCACCAGCTCCTCTATTCC	GGGATTATCGGCATTGTGAC
P7		
Interval	TTTTTGGTTTGCCCTTTTTG	GCTGCACGTGCGTAATTAAA
P8		
Interval	CCCGTCCACTGAAACTTCC	CCGCAGAGTCTGTTCCGTAT
P9	A CO A CTC OC A C A TCTTCTOC	A COTTOO CA CITTO A TOA CO A T
Interval	AGGACTGCCAGATGTTCTCG	AGCTGGGACTTGATCAGCAT
P10	CACCATAACTTCCCCCACAA	CAAACCCCACCAAAATCCTT
Interval P11	GAGGATAAGTTGGCCCACAA	GAAAGCCCAGCAAAATCCTT
Interval	TGCTGAACAGAACAGATCCA	TGGGGTACCGAAATTAACCA
P5'	IGEIGRACAGRACAGRICCA	TOOOTACCOAAATTAACCA
Interval	GAGGATAAGTTGGCCCACAA	GCGTCCGTGTTATCAGCTTT
P3'	0.100.11.11.01.1000001.11.1	
CG13442	TGCAATAACCTGAATTACGAGT	GTGTTTGTGCAAGGGCTCGC
RT PCR		
RP49 RT	AGCGCACCAAGCACTTCATC	GTGCGCTTGTTCGATCCGT
PCR		
CG13442	GAGTCCGCATCCAACGAATCGAT	GCGAGCCCTTGCACAAACAC

qRTPCR

Table 1: Primers

Alleles	P1	P2	P3	P4	P5	P6	P7	P8	P9	P10	P11	P5'	P3'
7A	-	-	+	-	-	-	+	-	-	-	-	+	-
18A	-	-	+	+	+	+	+	+	+	+	+	+	+
16A	-	-	-	-	+	+	-	-	-	+	+	-	+
19A	-	-	+	-	-	+	-	-	-	+	+	+	+
18B	-	-	+	-	-	+	+	-	-	+	+	+	+
WT(yw)	+	+	+	+	+	+	+	+	+	+	+	+	+

Table 2: Deletion Analysis of CG13442 mutants

+/- Refer to presence or absence of the band respectively

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