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Drosophila Inducer of MEiosis 4 (IME4) is required for Notch signaling during oogenesis

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N⁶-methyladenosine is a nonediting RNA modification found in mRNA of all eukaryotes, from yeast to humans. Although the functional significance of N⁶-methyladenosine is unknown, the *Inducer* of MEiosis 4 (IME4) gene of Saccharomyces cerevisiae, which encodes the enzyme that catalyzes this modification, is required for gametogenesis. Here we find that the Drosophila IME4 homolog, Dm ime4, is expressed in ovaries and testes, indicating an evolutionarily conserved function for this enzyme in gametogenesis. In contrast to yeast, but as in Arabidopsis, Dm ime4 is essential for viability. Lethality is rescued fully by a wild-type transgenic copy of Dm ime4 but not by introducing mutations shown to abrogate the catalytic activity of yeast Ime4, indicating functional conservation of the catalytic domain. The phenotypes of hypomorphic alleles of Dm ime4 that allow recovery of viable adults reveal critical functions for this gene in oogenesis. Ovarioles from Dm ime4 mutants have fused egg chambers with follicle-cell defects similar to those observed when Notch signaling is defective. Indeed, using a reporter for Notch activation, we find markedly reduced levels of Notch signaling in follicle cells of Dm ime4 mutants. This phenotype of Dm ime4 mutants is rescued by inducing expression of a constitutively activated form of Notch. Our study reveals the function of IME4 in a metazoan. In yeast, this enzyme is responsible for a crucial developmental decision, whereas in Drosophila it appears to target the conserved Notch signaling pathway, which regulates many vital aspects of metazoan development.

cell fate specification | meiosis | mRNA methylation | oocyte

N⁶-methyladenosine (N⁶-mA) is a modification found in less than 2% of all adenosines of eukaryotic pools of messenger RNA (1). Unlike other modifications, it does not change the identity of the base it modifies, and thus its presence cannot be inferred from the cDNA sequence. The existence of N⁶-mA in eukaryotic mRNA has been known for 4 decades, but its biological function has been elusive (2). However, the enzyme responsible for this modification is present, with very few exceptions, in all eukaryotes, implying a biological function for the N⁶-mA-modified mRNA species that has been kept throughout evolution.

Inducer of MEiosis 4 (IME4), the gene encoding N6-methyladenosine transferase in Saccharomyces cerevisiae, is required for entry into meiosis. It is expressed in mating-type locus a/mating-type locus α diploid cells and is required for mRNA accumulation of IME1, the transcription factor known to be the master regulator of yeast meiosis (3). Ectopic expression of IME4 in haploids is sufficient for these cells to initiate the meiotic program and attempt sporulation under starvation conditions, bypassing the requirement of mating-type heterozygosity (4). Although this mRNA modification in yeast is exclusively meiotic, and its dependency on Ime4 has been determined (5), the function of this modification in yeast meiosis remains unknown (6).

Our current knowledge of the biological function of *IME4* homologs in multicellular organisms is extremely limited, because mutants that compromise N⁶-adenosine methyltransferase activity have not been described in any metazoan. In *Arabidopsis thaliana*, the *IME4* homolog, *MTA*, has been shown to be an essential gene, but its role in development has not been explored (7). The existence of a *Drosophila* homolog of *IME4* (CG5933)

was reported in a comprehensive phylogenetic study of the evolution of methyltransferases (8). There exists a high degree of sequence similarity and conservation of the catalytic core among all eukaryotic homologs of *IME4* (5). Given the high degree of evolutionary conservation of *IME4* and its crucial function in the developmental decision of yeast to enter gametogenesis, we tested the hypothesis that *IME4* has a conserved function important for metazoan gametogenesis, focusing on oogenesis.

Drosophila oogenesis is a powerful system to identify gene functions controlling complex developmental events and signaling networks that are conserved in humans. The production of a fertilizable egg requires successive rounds of symmetry-breaking events that shape the follicle (egg chamber) and consequently define the future axes (polarity) of the embryo, as reviewed in ref. 9. Within the developmental unit of the fly ovary, the ovarioles, egg chambers composed of a germ-line cyst of 16 sister cells (15 nurse cells and one oocyte) encapsulated by a single layer of somatic follicle cells, emerge from the germarium and mature progressively. Egg-chamber maturation relies on continuous signaling between soma and germ line that drives morphogenetic processes and cell-fate determinations. Distinct follicle-cell fates are established early in oogenesis, specifically in germaria via Notch signaling, and are crucial for organizing the structure of the egg chamber, as reviewed in ref. 10.

Here we show that the *Drosophila* gene *Dm ime4* is essential for development, not solely for gametogenesis. We define a crucial role for the *Drosophila* gene in oogenesis. Our results indicate that Dm IME4 is required during oogenesis for processes that are regulated by Notch signaling and that involve somagerm-line interactions. We provide an experimental paradigm to investigate the plausible evolutionary conservation of IME4 in metazoans as a model for the function of the human gene.

Results

Dm ime4 Is Essential and Required for Fertility. Using the Model Organisms Best Hits search engine (11), we found that the *Dm ime4* predicted protein (CG5933) shares significant amino acid similarity with its homologs in *S. cerevisiae (IME4)*, *A. thaliana (MTA)*, *Mus musculus (METTL3)*, and *Homo sapiens (MT-A70)*, all of which are known or predicted to function as mRNA methyltransferases (Fig. S1A). The core catalytic domain is highly conserved among species, and its presence is a reliable predictor of enzymatic function (Fig. S1A) (5, 12).

To define the function of *Dm ime4*, we generated deletions by P-element excision mutagenesis (13). Deletions that removed the *Dm ime4* gene were homozygous lethal (Fig. 14). Two small deletions 5' to *Dm ime4* (Fig. S1B) reduced expression of the gene and were semilethal (we could recover viable adults, albeit at much lower than expected Mendelian frequencies) (Fig. 1B, Fig. S2, and Table S1). The homozygous animals died in larval and pupal stages (SI Materials and Methods). The rare adults

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The authors declare no conflict of interest.

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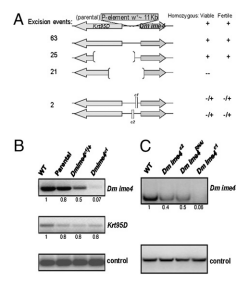


Fig. 1. Dm ime4 is essential for viability. (A) P-excision mutagenesis events that deleted the Dm ime4 ORF, partially or in its entirety, did not yield homozygous adult flies. Deletions of the adjacent Krt95D gene are viable. The small deletions 5' to the Dm ime4 gene greatly reduced female and male fertility. (B) The 5' deletion in the Dm ime4^{c1} (shown) specifically affects mRNA levels of Dm ime4 and does not affect mRNA levels of Krt95D. Transcript levels from whole flies were analyzed by RT-PCR in relation to a wild-type Oregon R control (numbers indicate relative abundance after quantification). Parental is the original P-element strain before P-element excision. Dm ime4^{c1}/+ is the heterozygous sibling balanced with TM3. Dm ime4^{c1} is the homozygous sibling rarely produced in the balanced stock. (C) Knockdown of Dm ime4 via RNAi. pUAST-based RNAi lines (VDRC) were used to knock down Dm ime4 using the ubiquitous act5C-GAL4 driver. mRNA from whole adult flies was analyzed as in B. The strongest phenotypes were observed with the homozygous Dm ime4c1 allele, which resulted in significantly lower levels of Dm ime4 than in the Dm ime4^{c2} allele or the Dm ime4^{RNAi} knockdown (Fig. S2).

recovered had significantly reduced fecundity. The severity of the semilethality and subfertility phenotypes correlated with decreased levels of Dm ime4 (Fig. 1 B and C and Fig. S2); i.e., the effects of the Dm ime4^{c1} allele were more severe than those of the *Dm ime4*^{c2} allele.

As an independent means to confirm the phenotype, we inhibited *Dm ime4* by RNA interference with transgenic lines obtained from the Vienna Drosophila RNAi Center (VDRC) that have no predicted or reported off-targets (14). By inducing double-stranded RNA homologous to Dm ime4 under actin 5c-GAL4 control to ablate gene function throughout development (Fig.1C and Fig. S2), the semilethality and reduced fertility phenotypes were reproduced. The effectiveness of reducing Dm ime4 levels via RNAi was comparable to the decrease in mRNA levels observed in *Dm ime4*^{c2} homozygous mutants (Fig. 1C and

To verify that the mutant phenotypes were caused by mutation of the Dm ime4 gene, we generated transgenic lines that expressed the Dm ime4 gene or a mutant variant using the GAL4- UAS inducible system for phenotypic complementation of Dm ime4c1 homozygous mutants. The semilethality of the *Dm ime4^{c1}* mutant was rescued by expressing the Dm ime4 gene under UAS control via actin 5C-GAL4, in that the recovery frequency of homozygous adults was equivalent to the expected Mendelian frequency for the cross (Table S1). In addition, the egg chamber phenotype described later in this report (see Fig. S6) also was rescued, and adults were fertile. Conversely, the semilethality of the Dm ime4^{c1} mutation was not rescued by expressing the Dm ime4 gene harboring two point mutations in the conserved catalytic core under UAS control via actin 5C-GAL, despite the presence of mutant protein levels comparable to the wild type (Fig. S3). The recovery

frequency of homozygous adults carrying the induced mutant transgene was equivalent to the frequency of the homozygous mutant without the induced transgene (Table S1). These point mutations are equivalent to the ones that result in loss of function in the S. cerevisiae homolog (5) and indicate conservation of the catalytic function of IME4 in Drosophila. In sum, our mutagenesis approach revealed an essential function for Dm ime4 and allowed the recovery of hypomorphic mutations that permitted us to investigate its role in adult flies.

Dm ime4 Is Expressed in Gonads. To test the hypothesis that Dmime4 is required for gametogenesis, we assessed its mRNA levels in ovaries and testes of adult flies. Dm ime4 was expressed in adult ovaries and testes but was not detectable in the carcasses from which gonads had been removed (Fig. 2 A and B). Our results were corroborated by cross-referencing the FlyAtlas gene expression database, where, as determined by microarray analysis, Dm ime4 transcript levels in the adult were highest in gonads and were undetectable elsewhere (15). By in situ hybridization to ovaries, Dm ime4 transcripts were detected in germaria, and hybridization signals persisted in later stages, when they were strongest in follicle cells (Fig. S4A). We chose to delineate the role of Dm IME4 in oogenesis because of the strong and specific expression of Dm ime4 in adult ovary. Moreover, we could recover homozygous mutant adults, albeit at very low frequencies, and these adults had significantly reduced fecundity. Furthermore, in Drosophila oogenesis the mechanisms of meiosis parallel those in most metazoans, and the developmental regulatory events are well defined.

 \mbox{Dm} IME4 Localizes to Soma and Germ-line Cells in Ovarioles. Wegenerated polyclonal antibodies against Dm IME4 to investigate protein levels and protein distribution in ovaries. Dm IME4 protein was detected in ovaries and not in protein extracts from carcasses after ovary removal (Fig. 2C). Consistent with reduced mRNA levels of Dm ime4 in Dm ime4^{c1} homozygotes, we observed significantly reduced levels of Dm IME4 protein compared with sibling heterozygous controls (Fig. 2D and Fig. S4B). By immunofluorescence, Dm IME4 was predominantly nuclear

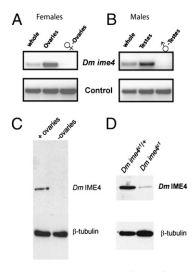


Fig. 2. Dm ime4 is expressed in the gonads of adult flies. mRNA from whole flies, dissected gonads (ovaries or testes, as indicated), or carcasses without gonads was analyzed as in Fig. 1. (A) Adult females. (B) Adult males. (C and D) Western blot using Dm IME4 antisera. (C) Protein extracts from wild-type Oregon R whole flies or their remains after ovary removal showing that Dm IME4 protein can be detected only in whole flies and not in those devoid of ovaries. (D) Protein extracts from ovaries isolated from females of the indicated genotype showing significantly reduced levels of Dm IME4 in homozygous mutant flies.

(Fig. 3 B–E), consistent with its function in posttranscriptional modification of mRNA and similar to the human homolog, MT-A70, which localizes to nuclei of HeLa cells (2). Dm IME4 was detected in somatic (prefollicle, follicle, and polar) cells and in germ line-derived cells (Fig. 3B). Dm IME4 protein levels were strong in the middle region of germaria (asterisks in Fig. 3B′), which is the region where 16-cell cysts of sister germ cells begin to be surrounded by a single layer of somatic follicle cells to form the emerging egg chamber (16). In later egg chambers throughout oogenesis, Dm IME4 protein was observed in all follicle cells but particularly strongly in polar cells (asterisks in Fig. 3C′ and D′). Dm IME4 protein also could be detected in the ooplasm (arrow in Fig. 3E′) and in the cells of the 16-cell cyst of early stages (arrow in Fig. 3B′). These patterns of expression indicated a likely role in egg chamber development, as addressed below.

Dm IME4 Is Required for Egg Chamber Development. Motivated by the protein-localization pattern of Dm IME4 throughout oogenesis in wild-type ovarioles and the decreased fecundity of Dm ime4 homozygous mutants, we predicted a role for Dm IME4 in oogenesis. To verify this prediction, we analyzed the morphology of ovarioles from homozygous mutants. The most significant and penetrant early phenotype was compound egg chambers with supernumerary nurse cells in the ovaries of the rare Dm ime4c1 homozygous mutant adults. This phenotype was observed in up to 70% of the mutant ovarioles examined (arrow in Fig. 4A' and Table S2). Mutant ovarioles that contained a significant number of compound egg chambers rarely matured past midoogenesis; egg chambers at later stages frequently were degenerating with pyknotic nuclei (asterisk in Fig. 4A'). Even in the absence of detectable compound egg chambers, seemingly normal ovarioles in the mutants also displayed a high frequency of degenerating egg chambers, and only 10% had mature egg chambers in vitellogenic stages (Table S2). Using an independent experimental approach, we also observed the compound egg chamber phenotype following heat-shock GAL4-induced *Dm ime4* RNAi in females (Fig. S5). Finally, the mutant phenotype is caused by the Dm ime4 mutation, because the Dm ime4 wild-type transgene can rescue the compound follicle phenotype of the mutants (Fig. S6).

We first addressed the nature of the germ cells in compound egg chambers. By labeling with the oocyte marker, Orb, we found that the mutant follicles contained two or more oocytes that failed to localize to the posterior end (arrowheads in Fig. 4B'). To determine whether the supernumerary germ cells originated from extra rounds of mitosis, we used an antibody that localizes to the cytoplasmic bridges (ring canals) between the oocyte and its sister nurse cells. In a normal 16-cell cyst there are four ring canals between the oocyte and its sister cells (Fig. 4C). We found that, with few exceptions (Fig. 4 C"), nearly all oocytes in compound follicles were surrounded by four ring canals; thus the majority of the cysts originated from the normal four rounds of mitotic divisions before entry of the oocyte into meiosis (Fig. 4C'and Table S2). In addition, the number of nurse cells in the compound follicles usually corresponded to a multiple of the number of oocytes detected; e.g., if three oocytes were detected, 45 nurse cells were present in that egg chamber.

We next investigated whether these oocytes in compound follicles entered meiosis by using an antibody against the central protein in the synaptonemal complex, C(3)G (Fig. 4 D and D'). We observed synaptonemal complex formation in these oocytes (Fig. 4 D'), indicating they entered meiosis.

Given that the compound follicles appeared to result primarily from the encapsulation of several 16-cell cysts into a common egg chamber, we examined follicle-cell morphology in the mutants. To trace the earliest signs of compound egg-chamber formation, we looked at the germaria of Dm ime4^{c1} mutants. A common feature of mutant germaria is the presence of two, sometimes more, partially encapsulated germ-line cysts located on the same plane (Fig. 5A'), instead of a single cyst file formed upon encapsulation as seen in the control germaria (Fig. 5A). In egg chambers that emerged from the germaria, we observed that follicle cells were disorganized and that specialized follicle-cell types, such as polar cells, frequently were absent or mislocalized (Fig. 5B'). Stalks (i. e., the cells that separate these follicles) were wider than normal stalks or were missing completely (Fig. 5 C and C'). In the most severe cases, there was a continuum of germ-line cysts from germaria to subsequent maturing stages (as discernable by the size of nurse cell nuclei; asterisk in Fig. 5A') with no apparent cyst individualization (arrow in Fig. 5A' and Table S2). Normally, Fasciclin III is down-regulated in epithelial follicle cells after

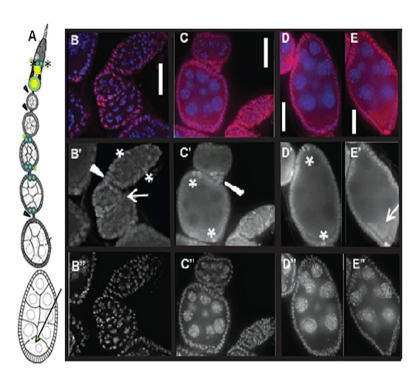


Fig. 3. Dm IME4 is abundant in follicle cells and also is detectable in early germ-line cysts. (A) Diagram of a Drosophila ovariole in the anterior-posterior orientation, with the germarium at the top. Asterisks at the middle of the germarium show the location of prefollicle cells (marked with asterisks in B'). Arrowheads show stalk cells connecting the egg chambers. Twin green dots marked with green asterisks show the location of anterior and posterior polar cells (marked with asterisks in C' and D'). Arrow shows the location of the ooplasm (arrow in E'). (B-E) Distribution of Dm IME4 protein (red) in ovarioles, showing localization in follicle cells and in germ-line cells. Panels are oriented so that top is anterior and bottom is posterior, as depicted in the cartoon in A. DAPI staining is blue. (B'-E') Dm IME4 single channel. (B''-E'') DAPI single channel. (B-B") Distribution of Dm IME4 protein in germaria showing prefollicle cells (asterisks in B'), stalk (arrowhead in B'), and early egg chambers (stages 2 and 3) where Dm IME4 is abundant in follicle and less abundant but reproducibly observed in germ-line cyst cells (arrow in B'). (C-C'') Germaria and stages 4 and 5, localization of Dm IME4 in stalk (arrowhead in C') and polar cells (asterisks in C'). (D-D'') Stage 7, localization of Dm IME4 in all follicle cells and abundantly in polar cells (asterisks in D') and forming a crescent shape (ooplasm). (E) Stage 8, localization of Dm IME4 in follicle cells, polar cells, and abundantly in ooplasm (arrow in E'). (Scale bars: 10 μ m.)

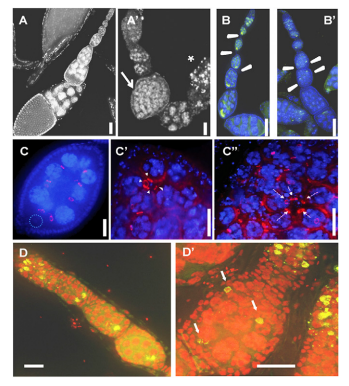


Fig. 4. Oogenesis defects in Dm ime4 mutants. All images are oriented with anterior at the top. (A, B, C, and D) Dm ime4^{c1}/+ control. (A', B', C', C'', and D') Dm ime4^{c1} homozygous. (A and A') DAPI nuclear staining showing a representative example of the control (A) and a Dm ime4^{c1} mutant ovariole (A') with a compound follicle (arrow in A') and a degenerating previtellogenic egg chamber (asterisk in A') (Refer to Table S2 for quantification of these phenotypes). (B and B') Compound follicles in Dm ime4^{c1} mutants have multiple oocytes [indicated by green Orb staining (arrowheads)] that fail to localize to the posterior end of the follicle. (C - C'') Most compound egg chambers of Dm ime4^{c1} mutants have multiple 16-cell germ-line cysts (15 nurse cells and one oocyte) from four rounds of mitosis, as indicated by the presence of four ring canals visualized by Kelch antibody (arrowheads in C') surrounded by a common layer of follicle cells. Dotted circle in C shows position of the posteriorly located oocyte. Five ring canals (arrows in C'') can be seen around the oocyte at low frequency. (D and D') Oocytes in Dm ime4^{c1} compound follicles entered meiosis and assembled the synaptonemal complex (arrows in D') as visualized by C(3)G antibody (green). DNA was detected with propidium iodide (red). (Scale bars: 10 µm.)

stage 2, remaining at high levels in later stages solely in polar cells (17). In contrast, this down-regulation of Fasciclin III was rarely observed in Dm imec1 mutant follicles (Fig. 5B'). Taken together, our results show severe defects in egg-chamber encapsulation and follicle-cell differentiation in Dm ime4 homozygous mutants that resembled defects in signaling between soma and germ line (18).

Notch Signaling Is Reduced in *Dm ime4* Mutant Follicle Cells. Defects in cyst encapsulation resulting in compound follicles and previtellogenic egg-chamber degeneration, the major phenotypes of Dm ime4 mutant ovaries, also are observed in mutants that affect Notch signaling: Delta, Notch itself, or Presenilin mutants (19). Therefore, we tested whether Notch signaling was compromised in Dm ime4 mutant ovarioles by using a transgenic line that permits visualization of Notch activation via a reporter fused to the Suppressor of Hairless regulatory region (20). The rare viable Dm ime4^{c1} homozygous adults with this reporter showed a significant decrease in the levels and in the number of follicle cells positive for Notch activation (Fig. 6 B and B') compared with heterozygous siblings (arrows in Fig. 6A and A'), implying that Dm IME4 is required for Notch signaling in oogenesis.

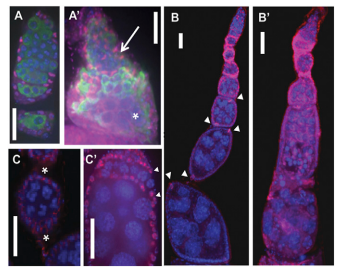


Fig. 5. Defects in cyst encapsulation and follicle-cell specification in Dm ime4 mutants. (A) Control germarium. (A') Representative germarium of a Dm ime4 homozygote in which cysts fail to follow an orderly pattern as they progress and have incomplete follicle-cell encapsulation (arrow). These trapped germ-line cysts can progress through other developmental stages, as shown by the increased size of the nurse cells at the bottom of this compound germarium (asterisk). Follicle cells were visualized with Traffic Jam (TJ, red). Germ-line cysts were visualized with VASA antibody staining (green). All nuclei were stained with DAPI (blue). (B and B') Polar cells indicated by arrowheads in control Dm ime4^{c1} /+ follicles (B) are aberrant or missing in $Dm ime4^{c1}$ compound follicles (B'). (C and C') Stalks seen in control Dm ime4^{c1} /+ (asterisks in C) frequently are missing in Dm ime4^{c1} homozygous mutants (C'), and egg chambers pack atop of each other. (Arrowheads in C' mark the developmental stage shown between the asterisks in C.) FasIII staining is, red; DAPI staining is blue. (Scale bars: 10 µm.)

Expression of Notch^{ICD} Rescues the Compound Egg Chamber Phenotype in Dm ime4 Mutant Ovaries. Because we observed a significant decrease in Notch signaling in the *Dm ime4* mutant follicle cells. we tested whether expression of a constitutively activated form of Notch (Notch CD, IntraCellular Domain) could suppress the ovarian phenotypes in *Dm ime4* mutants. Overexpression of Notch^{ICD} is deleterious before adulthood; thus Notch^{ICD} was induced under the control of the heat-shock promoter Hsp70 in females only after eclosion by performing daily heat-shock treatments for 5 d before dissection. An important aspect of eggchamber formation is that somatic follicle-cell differentiation is coordinated with germ-line events by cell-cell communication via Notch signaling. Disruption in soma-germ line communication leads to formation of aberrant egg chambers that frequently degenerate in previtellogenic stages and are not ovulated. As shown in Fig. 7, expression of Notch^{ICD} significantly reduced the frequency of compound egg-chamber formation in Dm ime4 mutant ovaries and allowed progression through oogenesis past the previtellogenic stages. The other significant phenotype of Dm ime4 mutant ovaries is the high frequency of degenerating egg chambers, but the suppression of this phenotype cannot be assessed accurately because of the high incidence of pyknotic nuclei that arise even in the controls during heat-shock treatment (Fig. 7). Of note, we observed significant suppression of the *Dm ime4* mutant phenotypes in the ovaries of females that carried the Notch^{ICD} transgene even in the absence of heat-shock treatment, probably because of leaky activity of the Hsp70 promoter at room temperature (21).

Discussion

Here we describe the role of the Dm IME4 mRNA N⁶-adenosine methyltransferase in the development of a metazoan organism. We find that, in contrast to the homologous gene in the unicel-

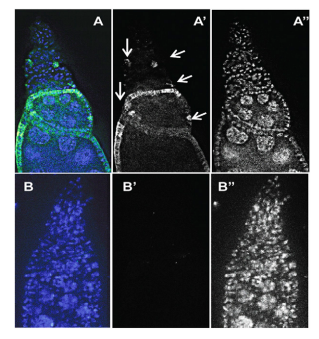


Fig. 6. Notch signaling is reduced in *Dm ime4* mutants. (*A* and *A'*) Notch signaling as detected in *Gbe+ Su(H)-lacZIFM7; Dm ime4^{C1}ITM6b* control (arrows indicate sites of Notch activity detected in the control). (*B* and *B'*) *Gbe+ Su(H)-lacZIFM7; Dm ime4^{C1}* mutants have significantly lower levels of Notch activity than the sibling *Gbe+ Su(H)-lacZIFM7; Dm ime4^{C1}ITM6b* controls as visualized by anti–β-galactosidase staining (green). *A* and *B* show merged anti–β-galactosidase (green) and DAPI (blue) channels; *A'* and *B'* show an anti–β-galactosidase (green) single channel. *A''* and *B''* show a DAPI single channel (white).

lular eukaryote *S. cerevisiae*, *Dm ime4* is an essential gene. In adults, we show that Dm IME4 is required for male and female fertility. In females, Dm IME4 is essential for oogenesis, and loss of function shows defects consistent with failure in soma–germ line interactions. We find that Notch signaling is reduced in *Dm ime4* mutants, suggesting a function for Dm IME4 in the Notch signaling pathway. Furthermore, Dm IME4 probably functions

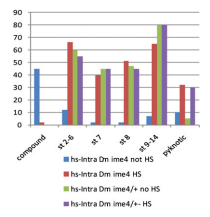


Fig. 7. Notch^{ICD} rescues egg-chamber defects in *Dm ime4* mutants. Bar graph shows quantification of 100 ovarioles for each of the genotypes and conditions assayed as described in the graph color key. Lines carrying the heat shock-inducible Notch^{ICD} construct are designated as hs-Intra, heat-shocked treated (HS), or not treated (no HS). Egg chambers in 100 ovarioles were scored and are represented on the *x* axis as follows: compound egg chambers (compound); previtellogenic stages 2–6 (st 2–6) and stage 7 (st 7); vitellogenic stage 8 (st 8) and stages 9–14 (st 9–14); and degenerating egg chambers (pyknotic).

upstream of this signaling pathway, because the expression of a constitutively activated form of Notch rescues the compound egg chamber phenotype of *Dm ime4* homozygous females.

The essential function of *Dm ime4* probably is a common feature in multicellular organisms. In A. thaliana, MTA, the Dm ime4 homolog, is essential for embryogenesis, because loss-offunction homozygous mutants are unable to proceed through embryogenesis past the globular stage and thus are unable to form differentiated tissues (7). Although the focus of this report is the function of Dm IME4 in oogenesis, homozygous mutant males also have reduced fertility; thus it will be interesting to determine the function of Dm ime4 in spermatogenesis and uncover similarities or differences in the roles of Dm IME4 in the ovary and testis. Further investigation of Dm IME4 function before its role in adult gametogenesis will reveal whether the protein controls cell differentiation in a variety of developmental contexts. Interestingly, the rare Dm ime4 mutant adults that are obtained have a high incidence of Notched wings, raising the possibility that Dm IME4 is required for Notch signaling in other developmental stages and tissues.

The defects in oogenesis that we observe when Dm ime4 function is compromised can be explained by failure of Notch signaling in follicle cells starting early in the germaria. Dm ime4 mutants and *Dm ime4* ablation via RNAi show defects in germ line-soma interactions leading to failure of follicle-cell differentiation, as shown by absence of stalks and polar cells and aberrant germ-line cyst encapsulation, similar to defects previously reported for Notch signaling mutants (19). Defects in somagerm-line communication, like those described for Notch signaling mutants, lead to the formation of aberrant egg chambers, which are eliminated via apoptosis (22). In addition to the phenotypic similarities between Dm ime4 and Notch signaling mutants, we observed significantly lower Notch reporter activity in Dm ime4 mutants than in sibling controls, indicating that Notch signaling is compromised by low levels of Dm IME4. Because the oogenesis phenotype of Dm ime4 homozygous mutants can be rescued fully by expressing an activated form of Notch, we show that Notch signaling is the pathway primarily affected in oogenesis in Dm ime4 mutants. Taken together, our data indicate that Dm IME4 is a key player in Notch signaling, probably functioning upstream of Notch activation. It will be interesting to determine how the enzymatic function of Dm IME4 affects this signaling pathway and whether transcripts harboring N⁶-mA-modified mRNA are involved in Notch signaling during soma-germ line interactions.

The function of yeast IME4 is to allow entry into meiosis; thus we expected a defect in meiotic entry in *Dm ime4* mutants. Because a complete deletion of *Dm ime4* is lethal, we could not investigate the phenotypic consequences of total absence of Dm IME4 protein in oogenesis. With this caveat, *Dm ime4* mutants that cause reduced fertility and ovary degeneration do not affect the onset of meiosis, as we can detect synaptonemal complex assembly in the oocytes of mutant egg chambers.

The protein expression of Dm IME4 and the phenotypes indicate a requirement in both the soma and the germ line. The mutant phenotype of compound egg chambers is caused by the inability of follicle cells to encapsulate a single 16-cell germ-line cyst, suggesting that the major role of Dm IME4 in oogenesis is in the somatic follicle cells. It is possible, however, that soma and germ line have different threshold requirements for Dm IME4 protein levels, and the reduction of Dm IME4 in the hypomorphic alleles may have affected the follicle cells primarily. We observed, albeit at low frequency, an extra round of mitotic germ-line cyst divisions. This phenotype is a consequence of Dm IME4 acting in the germ line, because it can be reproduced by RNAi knockdowns using germ-line drivers (Fig. S5). Taken together, our results suggest that Dm IME4 acts in both germ line and soma and plays a role in signaling between these two lineages during gametogenesis. In follicle cells this signaling appears to be accomplished via the Notch pathway.

In yeast, IME4 controls a crucial developmental decision in this unicellular eukaryote's life cycle: to continue mitosis or to enter the gametogenesis program (4). The present demonstration of developmental functions of *Drosophila* IME4 shows how a conserved function can be expanded in evolution, in this case for use in multiple developmental decisions and to target a signal transduction pathway that does not exist in yeast (23).

Materials and Methods

Ethics Statement. The protocol for production of polyclonal antibodies against Dm IME4 was reviewed and approved by the Committee for Animal Care of the Massachusetts Institute of Technology.

Strains and Genetic Manipulations. Bloomington strain 13927: y1; ry506 P{SUPor-P}KrT95DKG02054 harboring a P-element insertion marked with white+ in the intergenic region between Dm ime4 (CG5933) and Krueppel target at 95D (Krt95D) was used to induce P-element excision events via transposase to delete Dm ime4. The inducible Dm ime4 and catalytic domain mutant rescue constructs were generated by cloning the entire Dm ime4 cDNA in a pUASPbased Gateway vector as described (24), and their expression was induced by Actin5C-GAL4. Transgenic flies were provided by BestGene, Inc. The Gbe+ su(H)lacZ/FM7; TM2/TM6 transgenic line used to visualize Notch activity was provided by Sara Bray (University of Cambridge, Cambridge, UK) (25). The Hsp70-Notch^{ICD} transgenic line used to induce activated Notch by heat-shock treatment was a gift from Toby Lieber (Rockefeller University, New York, NY) (26).

Gene Expression. Transcript levels were quantified by RT-PCR. In situ hybridization was performed as previously described (27).

Antibody Generation and Protein Detection. Antigens and antibodies were generated as previously described (28). IME4 antigens were generated as GST-fusions of 0.8-kb and 1.2-kb fragments of Dm ime4 cDNA with high scores by antigenicity algorithm (29) and no predicted cross-reactivity. Anti-Dm IME4 antiserum was produced in guinea pigs (Covance, Inc). Western blots were performed as described (30).

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Immunofluorescence. Ovaries were dissected from 1- to 3-d-old females and fixed as previously described (31), with modifications given in SI Materials and Methods. All antibody dilutions were vol/vol. Dm IME4 antiserum from guinea pig was used at 1:1,000 dilution; mouse Fasciclin III (Developmental Studies Hybridoma Bank, DSHB) was used at 1:50; mouse Orb (DSHB) at 1:50 was amplified with anti-mouse biotin and visualized with FITC-streptavidin; mouse Kelch and mouse anti-Hts-RC (DSHB) at were used at 1:50; mouse C(3) G (a gift from Scott Hawley, Stowers Institute, Kansas City, MO) was used at 1:2,000, and samples for C(3)G detection were prepared as previously described (32). Guinea pig anti-Traffic Jam (a gift from Dorothea Godt, University of Toronto, Toronto, ON, Canada) was used at 1:5,000; rabbit anti-VASA (a gift from Ruth Lehmann, Skirball Institute, New York, NY) was used at 1:2,000; and anti- β -galactosidase (40-1a) (DSHB) was used at 1:50. Incubations with corresponding secondary fluorescent antibodies were done in PBS/3% BSA/0.1% Triton-X for 1 h at room temperature, at a dilution of 1:500. DAPI staining was done last, using a concentration of 1 µg/mL in PBS/ 0.5%BSA. Ovaries were mounted with VECTASHIELD (Vector Labs) or Aqua Poly/Mount (PolySciences) on polylysine-coated slides to immobilize ovaries for z-stack acquisition. Microscopic observations and imaging were done using a Nikon Eclipse Ti Epifluorescent inverted microscope with 10x and 20x dry objectives or 60x and 100x oil-immersion objectives, and images were analyzed using Nikon's NIS-Elements software.

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