

The 5'–3' exoribonuclease Pacman is required for normal male fertility and is dynamically localized in cytoplasmic particles in *Drosophila* testis cells

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The exoribonuclease Xrn1 is widely recognised as a key component in the 5'–3' RNA degradation pathway. This enzyme is highly conserved between yeast and humans and is known to be involved in RNA interference and degradation of microRNAs as well as RNA turnover. In yeast and human tissue culture cells, Xrn1 has been shown to be a component of P-bodies (processing bodies), dynamic cytoplasmic granules where RNA degradation can take place. In this paper we show for the first time that Pacman, the *Drosophila* homologue of Xrn1, is localized in cytoplasmic particles in *Drosophila* testis cells. These particles are present in both the mitotically dividing spermatogonia derived from primordial stem cells and in the transcriptionally active spermatocytes. Pacman is co-localized with the decapping

activator dDcp1 and the helicase Me31B (a Dhh1 homologue) in these particles, although this co-localization is not completely overlapping, suggesting that there are different compartments within these granules. Particles containing Pacman respond to stress and depletion of 5'–3' decay factors in the same way as yeast P-bodies, and therefore are likely to be sites of mRNA degradation or storage. Pacman is shown to be required for normal *Drosophila* spermatogenesis, suggesting that control of mRNA stability is crucial in the testis differentiation pathway.

Key words: P-bodies, RNA degradation, RNA stability, spermatogenesis, Xrn1.

INTRODUCTION

The control of RNA stability is a crucial element of gene regulation in all organisms. There is increasing evidence that specific, timed transcript degradation is critical for regulation of many cellular processes, including early development and RNA interference. The importance of RNA stability in developmental processes is revealed by the phenotypes of animals or plants where the expression of genes involved in RNA stability has been disrupted. For example, mice deficient in TTP (tristetraprolin), an RNA binding protein regulating stability of RNAs such as GM-CSF (granulocyte/macrophage colony-stimulating factor) and TNF (tumour necrosis factor), develop a systemic inflammatory syndrome with autoimmunity and myeloid hyperplasia (bone marrow overgrowth) [1]. In *Drosophila*, mutations in the 3'–5' deadenylase *ccr4* (*twin*) result in a level of sterility and maternal effect lethality in females [2]. Furthermore, *Caenorhabditis elegans* adults carrying mutations in *dicer* (*dcr-1*), a double-stranded endoribonuclease which plays a key role in the first steps of the RNA interference pathway, are sterile and have defects in developmental timing [3,4]. Finally, *Arabidopsis* mutants for DST1, which regulates the stability of RNAs such as CCL and SEN1, have defects in their circadian rhythm [5]. Therefore transcript degradation during development can be selective and also modulated, suggesting a little-studied layer of control of gene expression during development.

One of the key ribonucleases in the 5'–3' mRNA degradation pathway is the exoribonuclease Xrn1. This enzyme is known to be a processive 5'–3' exoribonuclease which degrades mRNAs after they have been de-capped [6–8]. Xrn1 is extremely well conserved in all eukaryotes, with homologues in humans (*xrn1*), *Drosophila* (*pacman*) and *C. elegans* (*xrn-1*) [9,10]. Indeed we

have previously shown that *Drosophila* Pacman has exonuclease activity *in vitro* and that it can complement a null XRN1 mutation in yeast [9–11]. Xrn1 is not only involved in the normal decay of mRNA but is required for the major pathway in nonsense-mediated decay [12], RNA interference [13] and degradation via microRNAs [14]. Mutations in *xrn1* in yeast or silencing of *xrn-1* in *C. elegans* lead to phenotypic defects, suggesting that this enzyme is required for normal cellular processes [6,8,9,15]. Our recent work has also shown that mutations in *pacman* in *Drosophila* result in defects in dorsal and thorax closure and reduced survival after wound healing [16].

Previous exciting studies have shown that 5'–3' degradation of cytoplasmic mRNAs seems to take place in specific particles known as P-bodies (processing bodies) [17–19]. In both yeast and human cells, P-bodies have been shown to harbour many enzymes of the basic decay machinery, including the decapping complex and the exoribonuclease Xrn1. Strong evidence that P-bodies are sites of normal mRNA decay, rather than merely storage sites for RNA decay factors, is that they accumulate mRNA decay intermediates [17,20], although mRNA degradation may also take place in sub-microscopic particles [21]. In mammalian cells, P-bodies are not only involved in normal mRNA turnover but have also been reported to be the sites of nonsense-mediated decay and miRNA (microRNA)-induced translational silencing [22,23]. P-bodies are dynamic structures that can vary in size and number according to stress conditions or the availability of untranslated RNAs [24,25]. Although P-bodies have been well studied in yeast and human tissue culture cells, they have rarely been observed in natural tissues. Their relevance to developmental processes is also unclear.

In this paper, we have examined the function of the exoribonuclease *pcm* (*pacman*) in *Drosophila* testis. We have

Abbreviations used: GFP, green fluorescent protein; P-bodies, processing bodies; TGF, transforming growth factor; UAS, upstream activator sequence; YFP, yellow fluorescent protein.

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chosen testis as a model system because they undergo a well defined differentiation pathway, remarkably similar to spermatogenesis in mammals [26]. In both flies and mammals, cells (spermatogonia) produced from self-renewing germ cells first mitotically divide to form spermatocytes and then, after a period of growth and transcription, shut off the transcriptional programme, enter meiotic divisions and differentiate into spermatids. Post-transcriptional pathways are highly regulated during the spermatogenesis process, with many transcripts being synthesised in primary spermatocytes and then translationally repressed until long after meiotic divisions [27]. The contribution of mRNA stability to spermatogenesis is as yet unclear.

This paper analyses the effect of the 5'–3' exoribonuclease *pacman* on male fertility in the fruit fly *Drosophila melanogaster*. We show that Pacman is required for normal testis development and male fertility. We also demonstrate that Pacman is located in cytoplasmic particles akin to P-bodies in both mitotically dividing spermatogonia and transcriptionally active spermatocytes. These particles also include the decapping activator dDcp1 and the helicase Me31B. These P-bodies, like those in human tissue culture cells, increase in size and number upon stress. Therefore we show, for the first time, that an exoribonuclease is crucial to normal male spermatogenesis.

EXPERIMENTAL

Drosophila stocks and mutant alleles

Fly stocks were cultivated on standard media at 25 °C in uncrowded conditions. The wild-type stock was Oregon R (Bloomington Stock Centre, Indiana University, IN, U.S.A.). The transgenic stock P{w+; YFP-dDcp1} (where YFP is yellow fluorescent protein) was provided by M.-D.L. [28]. The *pcm* mutants were generated by P-element excision and have been described previously [16]. Molecular characterization of the *pcm*³ and *pcm*⁵ alleles showed that the *pcm*³ allele comprises a deletion of 1522 bp, resulting in a frame-shift mutation. The resulting mutant protein is a C-terminal truncation of 197 amino acids relative to wild-type plus an extra 46 amino acids encoded from intergenic sequences; the 3' UTR was also deleted. The *pcm*⁵ allele is an intragenic deletion which removes 516 base pairs, resulting in a frame-shift. The *pcm*⁵ protein is C-terminally truncated by 348 amino acids compared to wild-type, with a 29 amino acid extension encoded from out-of-frame *pcm* sequences. The 3' UTR is intact in this allele. The wild-type protein is 1612 amino acids long.

Measurement of male fertility

The fertility of *pcm* mutant males was tested by crossing (at 25 °C) individual virgin males (control, *pcm*³ and *pcm*⁵) with individual wild-type virgin females, and 25 vials per strain were set up. The flies were left to mate for 2 days and then removed from the vials. F1 generation offspring were counted for 8 days after the first offspring hatched. The observed differences between the control and *pcm*³ and control and *pcm*⁵ were statistically significant in both cases (Mann-Whitney test, $W = 415.5$, $P < 0.0001$ and $W = 389$, $P < 0.0001$ respectively).

Measurement of sperm number

Seminal vesicles were dissected from 3-day-old virgin males ($n = 15$, at 25 °C) in a drop of PBS and transferred to a fresh drop of PBS. The seminal vesicles were then ruptured and the sperm inside allowed to dissociate. The preparations were left for 10 min and then allowed to dry. The time allowed for

dissociation and drying ensured that the sperm were well spread out. The slides were then fixed with 20 μ l of 4% paraformaldehyde solution in Hepes buffer and stained with DAPI (4',6-diamidino-2-phenylindole). Samples were mounted in 85% glycerol with 2.5% n-propyl gallate and cover slips were sealed with nail varnish. Sperm were then viewed at $\times 40$ with a Chroma long pass DAPI filter using a Leica fluorescent microscope. A video camera and monitor were used to view the slides and the sperm were counted. The observed differences between the control and *pcm*³ and also the control and *pcm*⁵ mutants were statistically significant in both cases (Mann-Whitney test, $W = 0$, $P < 0.0001$ and $W = 0$, $P < 0.0001$ respectively).

Measurements of testes

Testes were dissected from 2-day-old virgin males ($n = 20$, at 25 °C) in Ringer's solution (130 mM NaCl, 4.7 mM KCl, 1.9 mM CaCl₂ and 10 mM Hepes buffer, pH 6.9) and mounted in Ringer's solution on a microscope slide without a coverslip. The testes were visualized with light microscopy at $\times 20$ and the length and width (200 μ m from the tip of the testes) of the testes were measured with Zeiss AxioVision imaging system software. The observed differences for the length and width of the testes between the control and *pcm*³ and also the control and *pcm*⁵ were statistically significant in both cases (for the length of the testes: Mann-Whitney test, $W = 255$, $P < 0.025$ and $W = 361$, $P < 0.0001$ respectively; for the width of the testes: Mann-Whitney test, $W = 445$, $P < 0.0001$ and $W = 283$, $P < 0.0001$ respectively).

Western blotting

SDS/PAGE and Western blotting was performed essentially as described [29] with binding of antibodies being detected using an ECL Western blot reagent kit (Amersham). Primary polyclonal anti-PCM antibody (raised in rabbit) [16] was used at 1:2000. We are confident that this antibody (raised to the 54 kDa C-terminal portion of the protein) can detect the truncated mutant *pcm* proteins because it can pull down PCM from testis extracts (results not shown). The monoclonal anti-actin antibody (Sigma) was used at 1:10000 and primary antibody anti-dDcp1 (raised in rabbit), provided by Dr J. Wilhelm (Department of Embryology, Carnegie Institution of Washington, Baltimore, MD, U.S.A.), was used at a dilution of 1:2000. The secondary antibodies were monoclonal anti-rabbit HRP (horseradish peroxidase) conjugated antibody (Sigma) and monoclonal anti-mouse HRP conjugated antibody (Sigma), both used at 1:80000.

Immunofluorescence

For immunofluorescence, testes were dissected from 0–1-day-old males in Ringer's solution, with testes from approx. ten males being used per slide. The testes were transferred to a 20 μ l drop of Ringer's solution on a poly-L-lysine-treated slide and cut open with tungsten needles. Paraformaldehyde (20 μ l of 4% solution) in Hepes buffer was added and the testes were left to fix for 15 min at room temperature (20 °C). Testes were squashed by adding a cover slip, the slide was dipped in liquid nitrogen and the cover slip was removed with a scalpel. Testes were then stored in PBS+0.1% Tween-20 (PBS-T) until all the samples had been prepared. Samples were blocked for 30 min with PBS-T+10% FCS (fetal calf serum), then incubated with primary antibody diluted in blocking solution overnight at 4 °C. Testes were rinsed once then washed four times for 10 min each in PBS-T. Testes were incubated with secondary antibodies diluted in blocking

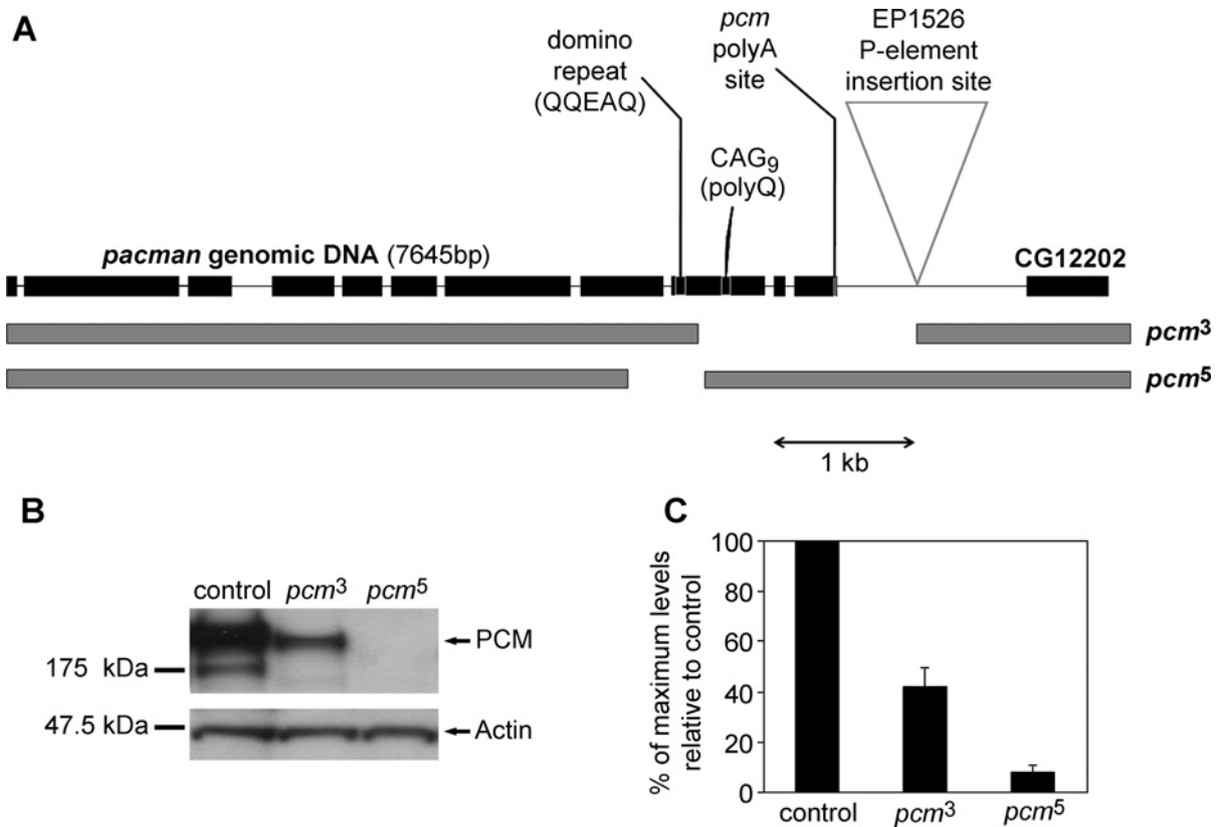


Figure 1 *pcm* and its mutants

(A) Diagram showing the genomic structure of *pcm*. Exons (■), introns (—), the 'domino' repeat (QQEAQ), the polyglutamine repeat (polyQ) and the poly-adenylation site is shown. The insertion site of the P-element used to generate deletions in the *pcm* gene is given, as is the downstream gene CG12202 (*Nat1*). The *pcm* alleles described in this paper are given below: gaps represent deletions at the genomic DNA level. (B) Western blotting of *pcm* hemizygous mutant male testes showing that the truncated *pcm* proteins are not detectable (*pcm*⁵) or are poorly expressed (*pcm*³). An actin loading control is shown below. (C) Histogram of normalized expression levels relative to loading controls. Means and standard errors are given.

solution for 2 h at room temperature. Samples were washed as before and mounted in 85 % glycerol with 2.5 % n-propyl gallate. Cover slips were sealed with nail varnish and the samples were imaged using a Leica TCS SP 2UV confocal microscope.

Primary antibodies used were rabbit anti-PCM, rabbit anti-dDcp1 (provided by J. Wilhelm) and mouse anti-Me31B (provided by Professor A. Nakamura, RIKEN Center for Developmental Biology, Kobe, Japan). For immunohistochemistry, the primary antibodies were used at a dilution of 1:500, 1:1000 and 1:1000 respectively. Secondary antibodies used were Cy-3-conjugated monoclonal mouse anti-rabbit IgG, Cy-5-conjugated goat anti-mouse IgG and FITC-conjugated goat anti-rabbit IgG (Jackson Immuno Research). The secondary antibodies were used at a dilution of 1:200.

The size of the cytoplasmic particles were measured using ImageJ software [30]. In each case the size of particles were measured for five different areas (35 μm \times 35 μm) to ensure repeatable results. The observed differences between the cases were statistically significant according to the Mann–Whitney test. The differences between the sizes of the cytoplasmic particles in *pcm*⁵ mutants under normal and stress conditions as detected using the dDcp1 antibody were rather low and the results of the Mann–Whitney test were inconclusive ($W = 236$, $P = 0.091$, which is > 0.05 but < 0.1); therefore we also used the exact permutation test. The percentage of co-localization was calculated using ImageJ software. In each case the percentage of co-localization was determined for three areas (25 μm \times 25 μm) to ensure repeatable results.

RESULTS

The 5'–3' exoribonuclease *pacman* affects male fertility in *Drosophila*

In order to study the effect of the 5'–3' exoribonuclease *pacman* on male fertility we examined males carrying mutations in the *pacman* gene. These mutations had been previously constructed using P-element excision of a P-element transposon located 584 nt downstream of the *pcm* polyA site, between *pcm* and the convergent gene *Nat1* on the X-chromosome at position 18D [16]. Two of the strongest alleles (*pcm*³ and *pcm*⁵) were used in this study (Figure 1A). Molecular characterization of these alleles showed that both encoded truncated proteins. The *pcm*³ allele was the result of a deletion from the P-element insertion site into the 3' end of the *pcm* gene and resulted in a C-terminal truncation of 197 amino acids relative to the wild-type. The *pcm*⁵ allele results from an intragenic deletion and encodes a protein C-terminally truncated by 348 amino acids (see the Experimental section for further details). Using a polyclonal antibody which specifically detects PCM [16] in Western blotting experiments, we showed that testes dissected from hemizygous *pcm*³ males produced truncated protein at levels significantly lower than wild-type, whereas *pcm*⁵ animals expressed very low levels of *pcm* protein (Figures 1B and 1C).

We next tested the effect of these *pcm* mutant alleles on male fertility by mating individual males to 3–5-day-old OregonR wild-type virgin females and counting the offspring. The *pcm*⁵ males were seriously compromised in their fertility, producing

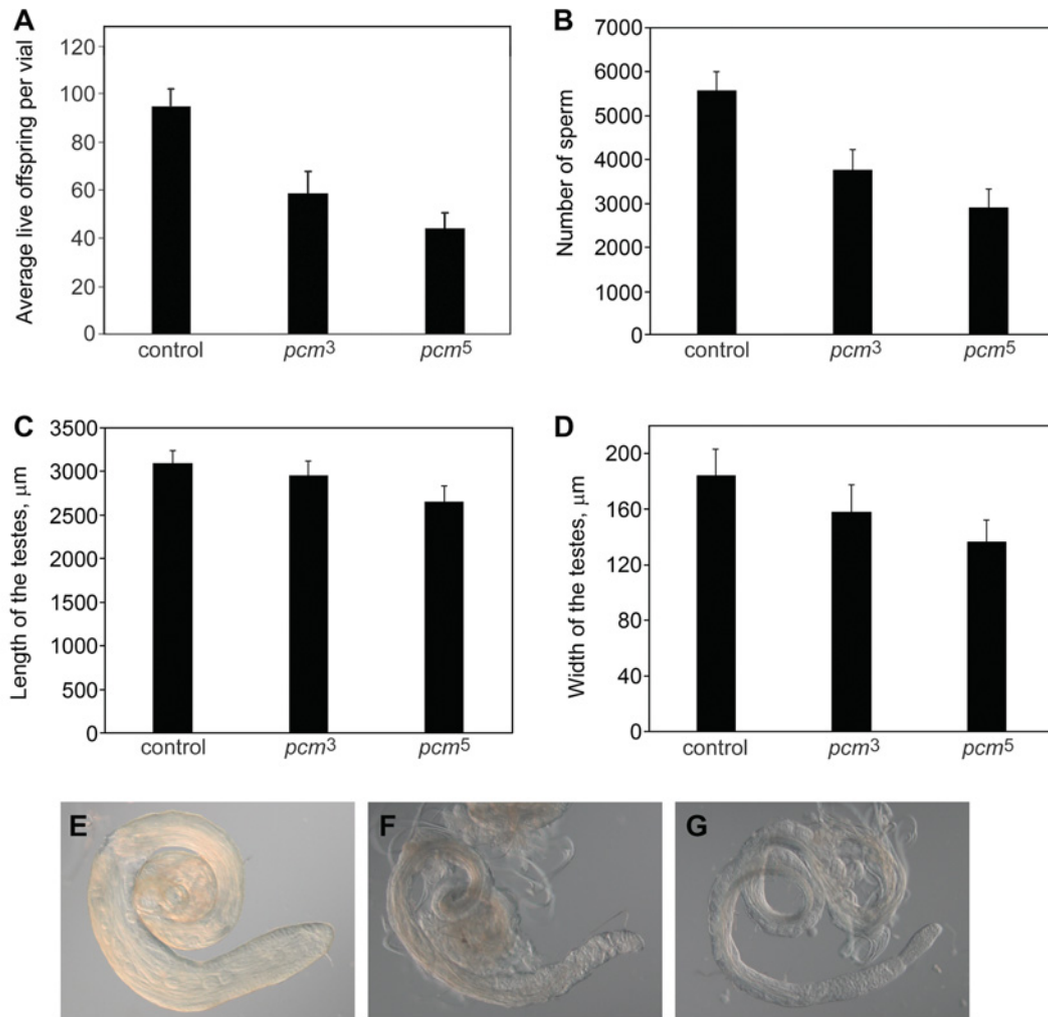


Figure 2 Effects of *pcm* on spermatogenesis

(A) *pcm* mutant males produce fewer offspring than controls. Means and standard errors are given. (B) *pcm* mutant males produce fewer mature sperm than controls. Seminal vesicles were dissected from 3-day old virgin males at 25 °C and the mature sperm counted. Means and standard errors are given. (C) and (D), *pcm* mutant testes are shorter (C) and thinner (D) than controls. Means and standard errors are given. (E–G) *pcm* mutant testes are smaller and 'weedier' than controls. (E), (F) and (G) represent testis dissected from 2-day-old wild-type, *pcm³* and *pcm⁵* males respectively.

only 47 % of the progeny of control males at 25 °C. (Figure 2A). Examination of testes from these mutant males showed that the mutant testes from 2-day-old *pcm⁵* males were much shorter and thinner than controls (Figures 2E–2G). Measurements of the length and width of these testes also showed that they were smaller than controls (Figures 2C and 2D). Male *pcm³* flies were intermediate in numbers of offspring and size of testes between *pcm⁵* and controls (Figure 2A, 2C and 2D). These smaller testes produced fewer sperm (68 % of wild-type in the case of *pcm³*; 52 % of wild-type for *pcm⁵*) (Figure 2B), which would account for the lower fertility of the *pcm* mutants.

Pacman is localized in cytoplasmic particles in wild-type testes

To start to understand how the exoribonuclease *pcm* might affect male fertility, we first examined the expression and localization of *pcm* protein in *Drosophila* testis. Using our anti-PCM antibody, we showed that PCM is well expressed at the tip of the testis, in the spermatogonia where the initial mitotic divisions take place (Figure 3A) [26]. Closer examination revealed that PCM is

located in foci in these cells (Figure 3B). To confirm that PCM is located in cytoplasmic granules in living cells, we generated a UAS-GFP-PCM transgenic that expresses GFP (green fluorescent protein) fused in-frame to the N-terminus of PCM under the control of the UAS (upstream activator sequence) enhancer. When this GFP-PCM is expressed in testes using the *bam*-GAL4 driver, GFP-PCM also accumulates in particles (Figure 3C). However, in *bam*-Gal4 UAS-GFP-PCM flies, the GFP-PCM expression also appears in the fusome, a membranous organelle that allows endoplasmic reticulum connectivity between cells [31]. This is most likely to be because it is over-expressed from the foreign promoter. The expression of PCM in spermatogonia suggests that it is required for the amplification or maintenance of these dividing spermatogonia prior to their entry into pre-meiotic S-phase as primary spermatocytes.

We then examined the localization of PCM in primary spermatocytes, which are cells that do not divide but undergo massive transcription and growth. In these cells, PCM is located in discrete, cytoplasmic foci, as detected in both GFP-PCM flies (Figure 3D) and immunocytochemistry experiments (Figure 3E). The staining

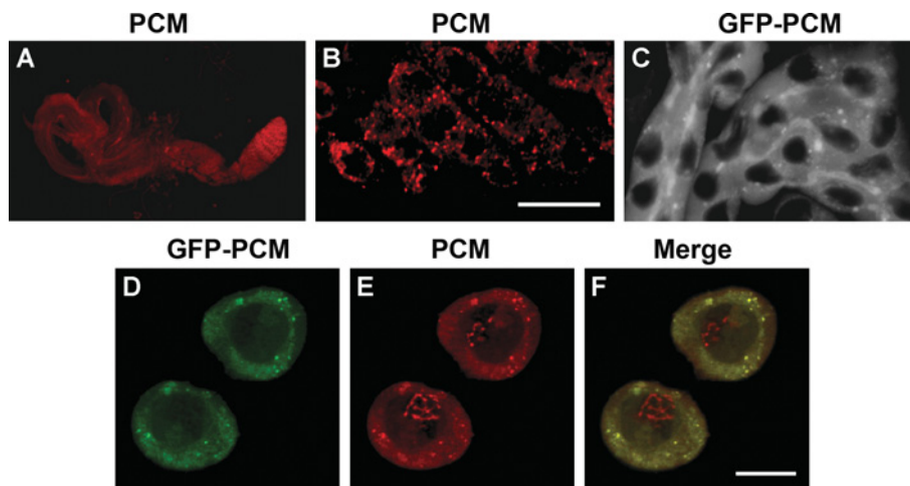


Figure 3 Localization of PCM in testis cells

(A) PCM antibody staining shows that PCM is highly expressed at the tip of the testis. (B) PCM is expressed in cytoplasmic foci in spermatogonia. Scale bar: 10 μ m. (C) GFP-PCM is expressed in cytoplasmic foci in spermatogonia from transgenic males. (D–F) GFP-PCM is expressed in spermatocytes in transgenic males (D) and co-localizes with PCM detected using PCM antibody (E). (F) Merged image. Scale bar: 10 μ m.

of granules in testis from transgenic flies expressing GFP-PCM co-localizes with PCM antibody staining (Figure 3F). These experiments show that PCM is also required at this stage of differentiation.

This is the first time that cytoplasmic foci that may be related to P-bodies have been seen in *Drosophila* testis. These particles are reminiscent of Xrn1 particles in mouse cells under conditions where the cells are not stressed, and native levels of protein (rather than protein expressed from a foreign promoter) are observed. In human tissue culture cells such as HEK (human embryonic kidney)-293 cells, P-bodies can also be present in unstressed cells [24]. This result shows that PCM is dynamically localized in testis cells and suggests that these particles could be *Drosophila* P-bodies.

Pacman is co-localized with dDcp1 and Me31B in testis cells

In both yeast and human tissue culture cells, P-bodies have been shown to contain enzymes required for decapping of mRNA (dDcp1 and dDcp2) as well as the helicase Me31B/Dhh1/p54 [17–20,32,33]. In both human and yeast cells, Dcp1 is generally regarded as a marker for P-bodies [34]. To determine whether the PCM foci seen in testis cells might be P-bodies, we investigated whether dDcp1 is located in cytoplasmic granules and if PCM is co-localized with dDcp1. Use of both an anti-dDcp1 antibody and a dDcp1-YFP transgenic show that dDcp1 is localized in cytoplasmic granules in both spermatogonia and primary spermatocytes (Figures 4A and 4B). Co-localization experiments showed that dDcp1 is largely, but not completely, co-localized with PCM in spermatocytes in that 95% of PCM protein visualized is co-localized with dDcp1 (Figures 4C–4E).

Another protein known to be critically important in P-body formation in *Saccharomyces cerevisiae* is the RNA helicase Dhh1. This enzyme is highly conserved, with the known homologue in *Drosophila* being Me31B [35]. Using an antibodies to Me31B, dDcp1 and PCM we showed that Me31B co-localizes with dDcp1 (Figures 4I–4K) and Me31B co-localizes with PCM (Figure 4F–4H) in spermatocytes (95% in both cases). Close examination of these particles revealed that co-localization of Me31B and dDcp1 was almost complete, whereas both of these were partially

co-localized with PCM (see insets in Figures 4E, 4H and 4K). This suggests a structure to these particles which has not been previously observed. The co-localization between PCM, dDcp1 and Me31B strongly suggest that these particles are akin to P-bodies.

Pacman mutations affect the size and numbers of testis P-bodies

In the yeast *S. cerevisiae*, cells mutant for Xrn1 are blocked in 5'–3' exonuclease degradation (*xrn1* Δ), and show dramatic increases in size and number of P-bodies [17]. If these testis granules are the sites where mRNA degradation takes place, we would therefore expect the size of P-bodies to increase when the Xrn1 homologue Pacman is depleted. To test this prediction we first immunostained hand-dissected testes using our PCM antibody. In *pcm*⁵ mutants, only diffuse, low intensity staining was seen with no cytoplasmic foci visible (Figure 5C). This result is consistent with our Western blotting experiments (Figure 1B) where no PCM band was visible. The *pcm*³ mutants showed diffuse, faint staining, with few obvious cytoplasmic particles (Figure 5B). Note that this staining in Figures 5(B) and 5(C) was visualized using a higher power of the laser than the other panels. Therefore, as expected, these hypomorphic mutations result in a reduction of staining in these mutants. In addition, in the case of the *pcm* mutation, which encodes a C-terminal truncation of PCM (Figure 1A), these experiments also suggest that the C-terminus of PCM may be important in localizing PCM to P-bodies. The C-terminus of PCM includes a polyglutamine repeat, which is deleted at the protein level in the *pcm*³ mutant. These data suggest that protein motifs within the C-terminus of PCM, possibly including the polyglutamine repeat, are involved in the localization of PCM to P-bodies.

We then examined the effects of these *pcm* mutations on P-bodies using the P-body marker dDcp1. In these *pcm* mutants, P-bodies are dramatically increased in size and number in a similar way to that observed in yeast cells mutant for Xrn1 (Figures 5D–5F). Quantification of the size difference reveals that the particles are increased 2.7 times in *pcm*⁵ mutants compared with controls (Figure 5G). These results provide additional evidence that the testis granules marked by PCM and Dcp1 are equivalent to yeast P-bodies.

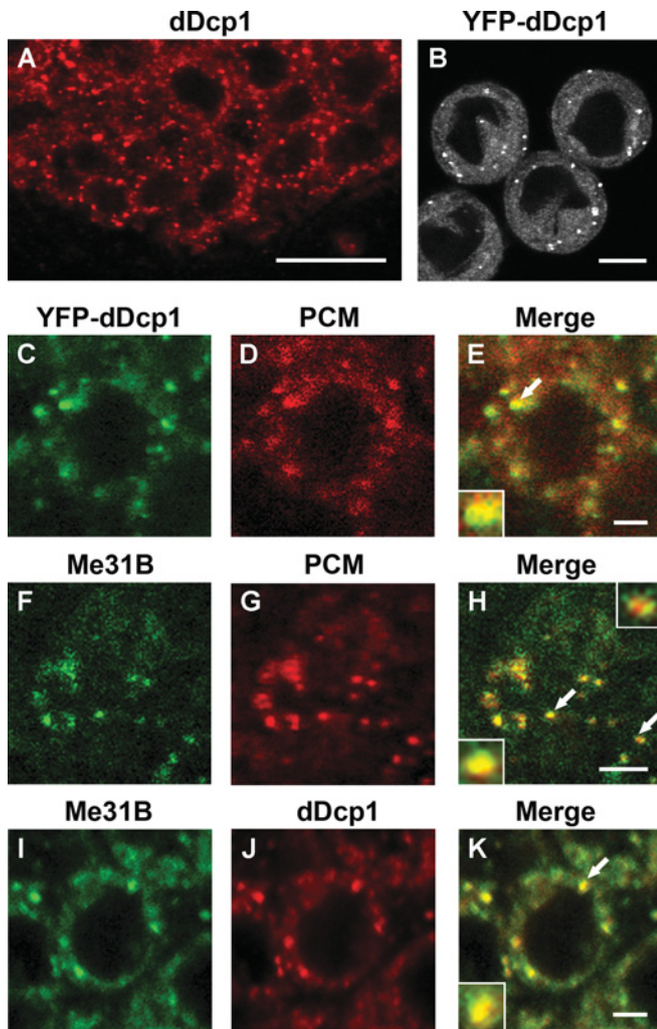


Figure 4 PCM is partially co-localized with the mRNA decapping enzyme dDcp1 and the helicase Me31B in P-bodies

(A) and (B) dDcp1 antibody staining (A) and YFP-dDcp1 expressed in testis cells in transgenic males (B) showing that dDcp1 is expressed in cytoplasmic foci in testis cells. Scale bars: 10 μm . (C–E) YFP-dDcp1 expressed in testis cells in transgenic males (C) is partially co-localized with PCM detected using PCM antibody (D). (E) Merged image. Scale bar: 2 μm . (F–H) The helicase Me31B (F) is partially co-localized with PCM (G) in cytoplasmic particles in testis cells. (H) Merged image. Scale bar: 2 μm . (I–K) The helicase Me31B (I) is partially co-localized with dDcp1 (J) in cytoplasmic particles in testis cells. (K) Merged image. Scale bar: 2 μm .

The increase in dDcp1 localization to P-bodies could be due to an increase in dDcp1 expression or recruitment of dDcp1 from the cytoplasm to these RNA granules. To test this, we performed Western blotting experiments on hand-dissected testis from wild-type and *pcm* mutants using the dDcp1 antibody. These results showed that the amount of dDcp1 decreases in *pcm* mutants (Figures 5H and 5I), supporting the notion that the increase in Dcp1 staining in P-bodies in *pcm* mutants is due to massive recruitment of this protein to P-bodies.

Pacman foci increase in size and number upon heat stress

In yeast and human tissue culture cells, environmental stresses can result in increases in sizes and numbers of P-bodies. Stresses such as glucose deprivation, osmotic stress and UV irradiation increase the size and numbers of P-bodies in yeast cells [24]. Although heat stress has not been reported to lead to an increase in size

or numbers of P-bodies in yeast [25], the heat stress used was relatively mild (37 °C), so may not have been strong enough to elicit a P-body response [36]. In human DU145 cells, heat shock (44 °C for 1 h) can result in an increase in numbers of P-bodies [24].

If these testis granules are akin to human P-bodies, we would expect them to increase in size when the male flies are stressed. To test this prediction we subjected male *Drosophila* to heat stress. We used heat stress as it is easy to administer to live flies and has already been shown to affect male fertility without substantially compromising viability [37]. Our experiments show that, in testes dissected from males subjected to heat shock (37 °C for 1 h), the numbers and sizes of granules marked by PCM increases 4.1-fold (Figures 6A–6C). In addition, granules marked by dDcp1 also increase between 1.3- and 2.1-fold in wild-type and mutant testis when males are heat-shocked (Figures 6D–6L). The response of these testis granules to heat stress is very similar to that seen in human tissue culture cells, confirming that these granules are likely to be P-bodies [24]. These results raise the possibility that P-bodies are important in the response of testis cells to stress and suggest that mRNA turnover and its relationship to translation is similar between *Drosophila* and humans.

DISCUSSION

Pacman is located in particles akin to P-bodies in *Drosophila* testes

PCM is located in cytoplasmic particules in both spermatogonia and spermatocytes. What is the function of these particles? Results from the present study suggests that they are akin to P-bodies, which are thought to be involved in the degradation of RNAs. The evidence for this is that PCM (an exoribonuclease) is co-localized with the decapping activator Dcp1 and the helicase Me31B (a Dhh1 homologue). In addition, we have shown that PCM is physically associated with dDcp1. Dcp1 is often regarded as a marker for P-bodies [34]. In addition, these particles behave in the same way as yeast and human P-bodies in that they increase in size upon stress (Figure 6), and also in mutants where members of the 5'–3' pathway are downregulated.

Although elegant work in yeast has shown that P-bodies accumulate partially degraded mRNAs, further work has shown that P-bodies can also store mRNAs. Therefore the function of P-bodies in yeast is not yet clear. However, our recent work examining PCM localization of PCM in *Drosophila* egg chambers has shown the existence of two types of particle containing PCM, dDcp1 and dDcp2: one type in the nurse cell cytoplasm which responds to stress and depletion of RNA decay factors and the other type in the oocyte, which is resistant to these factors [38]. Since the oocyte particles are also co-localized with known translational repressors such as Oskar, these appear to be storage bodies, rather than particles involved in mRNA degradation [38]. Furthermore, in *Drosophila* neurones, PCM is co-localized with translation repression factors such as dFMR1, Cup and Pumilio in neuronal granules [39], suggesting that it is also involved in translational repression in these tissues. Although we cannot rule out that the PCM particles we see in testis are storage granules, their response to stresses and depletion of mRNA factors, together with their small size, suggests to us that they are involved in RNA degradation.

The large size of spermatocytes has allowed us to analyse the internal structure of PCM-containing granules in a natural tissue in a way that has not previously been possible in yeast or human tissue culture cells. As shown in Figure 4, Me31B and dDcp1 are almost completely co-localized with each other in these particles, whereas PCM is partially co-localized with both. Note that this

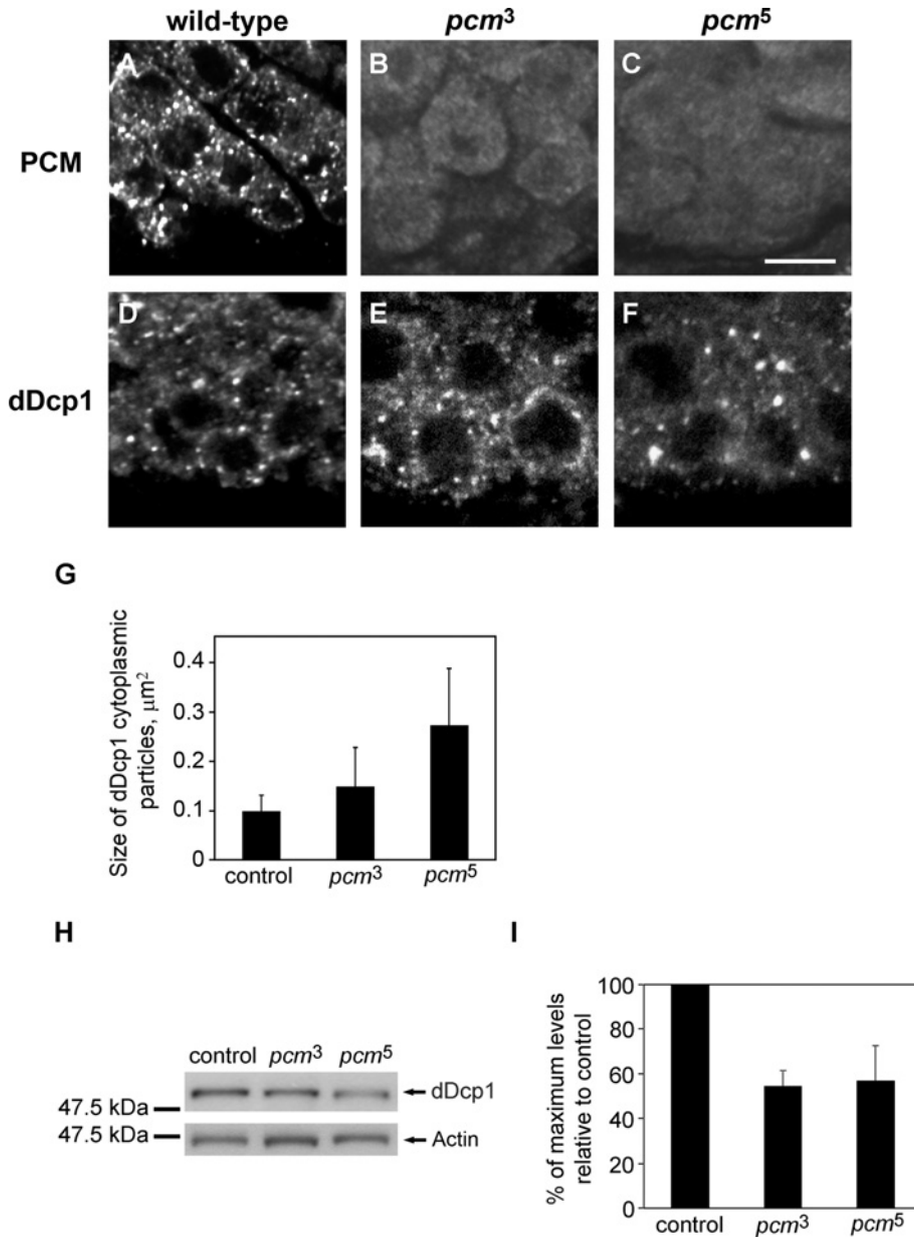


Figure 5 Effect of *pcm* mutants on cytoplasmic particle size and dDcp1 expression

(A–C) PCM antibody staining shows that cytoplasmic particles are more diffuse in *pcm* mutants than controls. (A), (B) and (C) represent wild-type, *pcm³*, and *pcm⁵* testis respectively. Images for *pcm* mutants in (B) and (C) were taken at a higher power of confocal laser since no signal is seen when a power equivalent to panel (A) was used. Scale bar: 5 μm . (D–F) dDcp1 antibody staining shows that cytoplasmic particles are larger in *pcm* mutants than control. (D), (E) and (F) represent wild-type, *pcm³* and *pcm⁵* testis, respectively. (G) dDcp1 cytoplasmic particles are larger in *pcm* mutants than control. Means and standard errors are given. The observed differences for the size of the particles between the control and *pcm³* and control and *pcm⁵* were statistically significant in both cases (Mann-Whitney test, $W = 422.5$, $P = 0.0106$ and $W = 632.5$, $P < 0.0001$ respectively). (H) Western blotting of *pcm* hemizygous mutants (*pcm³/Y* and *pcm⁵/Y*) showing that dDcp1 is expressed at lower levels in mutants compared to controls. An actin loading control is shown below. (I) Histogram of normalized expression levels relative to loading controls. Means and standard errors are given.

co-localization has been seen using antibody staining of wild-type testis, showing that co-localization is not an artifact arising from over-expression of fusion proteins. These results are similar to that seen in the cytoplasm of nurse cells in *Drosophila* egg chambers, where PCM is partially co-localized in particles with Me31B and dDcp1 [38,39]. These data suggest a scenario where the helicase Me31B and the decapping activator dDcp1 bind mRNA together and then bring it to the exoribonuclease PCM. Further experiments using GFP- and YFP-tagged proteins in live testis imaging are in progress to understand the dynamic nature of these particles in these natural tissues.

Pacman affects male fertility

Our results from the present study show that the exoribonuclease *pcm* is required for normal male fertility in *Drosophila melanogaster*. Examination of *Drosophila* males carrying hypomorphic mutations in *pcm* show that their testes are small and abnormal and that they produce fewer offspring compared to controls. The expression levels of *pcm* are correlated with the number of offspring produced, as *pcm⁵* males (which express undetectable levels of truncated protein) produce fewer offspring than *pcm³* males (expressing higher levels of mutant protein). These results

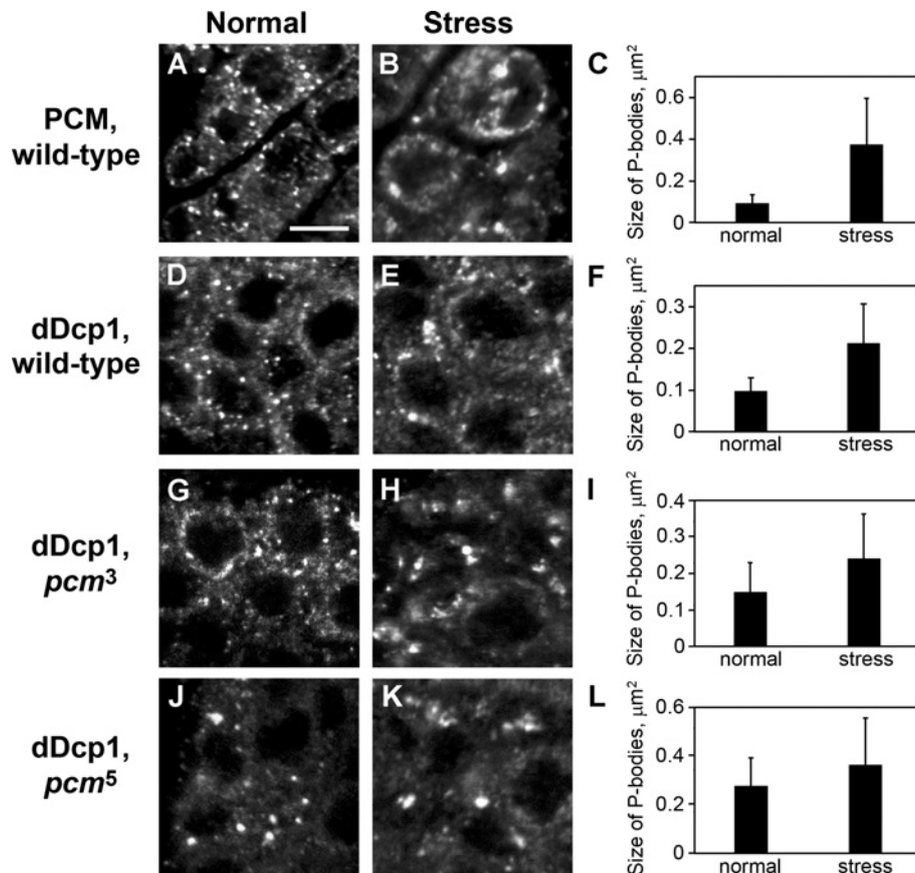


Figure 6 Under stress conditions, the cytoplasmic particles containing PCM and dDcp1 increase in size

(A–C) Particles containing PCM in wild-type spermatogonia increase in size upon stress. (A) Normal condition. (B) Stress condition. (C) PCM cytoplasmic particles increase in size under stress conditions. Means and standard errors are given. The observed difference between the normal and stress conditions was statistically significant (Mann-Whitney test, $W = 2.5$, $P < 0.0001$). (D–L) Particles containing dDcp1 in wild-type (D–F), pcm^3 (G–I), and pcm^5 (J–L) spermatogonia increase in size upon stress. (D, G and J) Normal condition. (E, H and K) Stress condition. Scale bar: 5 μm. (F, I and L) dDcp1 cytoplasmic particles in wild-type (F), pcm^3 (I), and pcm^5 (L) increase in size upon stress. Means and standard errors are given. The observed difference between the normal and stress conditions in wild-type, pcm^3 and pcm^5 was statistically significant (wild-type, Mann-Whitney test, $W = 49$, $P < 0.0001$; pcm^3 , Mann-Whitney test, $W = 125.5$, $P = 0.0054$; pcm^5 , exact permutation test, $T = 262$, $P = 0.049$).

show for the first time that an exoribonuclease is required for spermatogenesis.

How could an exoribonuclease, which is relatively non-specific *in vitro* [11], affect spermatogenesis? PCM is expressed at the tip of the testis, in the spermatogonia, showing that PCM is required in these mitotically-dividing cells. In these cells, it is possible that *pcm* perturbs TGF (transforming growth factor)- β signalling, as the mutant testes strongly resemble those mutant in the TGF- β ligand Punt [40]. Indeed, in complementary experiments, we have shown that *pcm* genetically interacts with *puckered*, a member of the JNK (c-Jun N-terminal kinase) signalling pathway that can regulate TGF- β signalling [41]. TGF- β signalling is required for maintenance of the germline stem cells: lack of the TGF-ligand Gbb (Glass bottom boat) results in a reduction of stem cells with the concomitant reduction of mitotically-dividing spermatogonia. In spermatocytes, PCM may disrupt differentiation of spermatocytes by the inappropriate upregulation of target RNAs. Since *pcm* is highly conserved, and the spermatogenesis process has similarities between *Drosophila* and humans, it is also likely that human Xrn1 affects human spermatogenesis. Our further work to elucidate the targets of PCM in *Drosophila* spermatogenesis will therefore promote understanding of gene regulation in human germ cells.

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