

# Regulation of transcription of meiotic cell cycle and terminal differentiation genes by the testis-specific Zn-finger protein *matotopetli*

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## Summary

A robust developmentally regulated and cell type specific transcriptional programme is activated in primary spermatocytes in preparation for differentiation of the male gametes during spermatogenesis. Work in *Drosophila* is beginning to reveal the genetic networks that regulate this gene expression. The *Drosophila* *aly*-class meiotic arrest loci are essential for activation of transcription of many differentiation-specific genes, as well as several genes important for meiotic cell cycle progression, thus linking meiotic cell cycle progression to cellular differentiation during spermatogenesis. The three previously described *aly*-class proteins (*aly*, *comr* and *achi/vis*) form a complex and are associated with chromatin in primary spermatocytes. We identify, clone and characterize a new *aly*-class meiotic arrest gene, *matotopetli* (*topi*), which

encodes a testis-specific Zn-finger protein that physically interacts with *Comr*. The *topi* mutant phenotype is most like *achi/vis* in that *topi* function is not required for the nuclear localization of *Aly* or *Comr*, but is required for their accumulation on chromatin. Most target genes in the transcriptional programme depend on both *topi* and *achi/vis*; however, a small subset of target genes are differentially sensitive to loss of *topi* or *achi/vis*, suggesting that these *aly*-class predicted DNA binding proteins can act independently in some contexts.

Supplemental data available

Key words: Spermatogenesis, Transcription, Chromatin, Meiosis, Differentiation.

## Introduction

Tissue and cell type specific transcriptional activation is a fundamental feature of cellular differentiation that mediates many changes in cell behaviour and morphology during development. One of the most astonishing developmentally regulated changes in cell morphology occurs in the production of sperm. The cellular differentiation events that constitute spermiogenesis require many gene products used at no other time in development. The unique features of male gamete differentiation are reflected in a developmentally specific transcription programme that initiates in primary spermatocytes in preparation for spermiogenesis. In *Drosophila* spermatogenesis, transcription is essentially shut down as mature primary spermatocytes enter the meiotic divisions. Therefore, gene products required during differentiation are largely produced from transcripts expressed pre-meiotically (Olivieri and Olivieri, 1965; Schafer et al., 1995). A two-part genetic module, the meiotic arrest genes, is responsible for most of this transcriptional activation, and ensures that a large number of genes required for normal

cellular morphogenesis are co-expressed in a cell type and developmental stage-specific manner (Lin et al., 1996; White-Cooper et al., 1998).

Males mutant for any of the meiotic arrest genes [including *always early* (*aly*), *cannonball* (*can*), *meiosis I arrest* (*mia*), *spermatocyte arrest* (*sa*), *cookie monster* (*comr*), *achintya/vismay* (*achi/vis*) and *no-hitter* (*nht*)] are viable but sterile. Testes from flies mutant for any one of these genes contain morphologically normal spermatogenic cells up to mature primary spermatocytes. However, the mutant cells then arrest and fail to initiate either the meiotic divisions or spermatid differentiation (Ayyar et al., 2003; Jiang and White-Cooper, 2003; Lin et al., 1996). Although transcription of a number of broadly expressed genes (such as *cyclin A*) is normal in the mutant spermatocytes, transcription of many spermiogenesis genes (e.g. the mitochondrial fusion protein *fzo*) is very low or undetectable. The *aly*-class meiotic arrest genes (*aly*, *comr*, *achi/vis*) are likely to act in a pathway distinct from the *can*-class genes, because they are required for transcription of a wider range of target genes. In addition to a

role in transcription of spermiogenesis genes, *aly*-class genes are also essential for the transcriptional activation in primary spermatocytes of several genes required for progression into the meiotic divisions (namely *twine*, *cycB*, *boule*). By contrast *can*, *mia*, *sa* and *nht* (*can*-class genes) are not required for transcription of these cell cycle genes; however, they are required for translation of *twine*. The *can*-class meiotic arrest genes analysed to date encode testis specific homologues of more generally transcribed TATA-binding protein associated factors (TAFs) (Aoyagi and Wassarman, 2000; Hiller et al., 2001) (M.H. and M.T.F., unpublished). TAFs are subunits of the basal transcription factor TFIID, which is involved in recruitment of the RNA polymerase II holoenzyme to the proximal promoter region. Thus, the *can*-class genes are likely to activate full levels of transcription of spermatid differentiation genes through intimate association with target promoters (Hochheimer and Tjian, 2003).

Not all genes transcribed in primary spermatocytes are controlled by the meiotic arrest pathway, so it is likely the targeting of particular promoters is mediated through sequence-specific DNA-binding activity. Thus, we would expect that one (or more) of the meiotic arrest genes encode a DNA-binding protein(s), whereas others are likely to encode regulatory proteins. *aly* is one of two *Drosophila* homologues of the *C. elegans* vulval differentiation regulator, *lin-9* (Beitel et al., 2000; White-Cooper et al., 2000). The SynMuvB pathway, in which *lin-9* acts, is thought to negatively regulate adoption of vulval fate and promote hypodermal fate, through activation of a histone deacetylase chromatin remodelling complex (Ferguson and Horvitz, 1989; Lu and Horvitz, 1998; Solari and Ahringer, 2000). *comr* encodes a novel acidic protein with no significant similarities to other proteins in current sequence databases (Jiang and White-Cooper, 2003). *achi/vis* was the first meiotic arrest locus to be shown to encode products with sequence-specific DNA-binding activity (Ayyar et al., 2003; Wang and Mann, 2003). The TALE class homeodomain proteins encoded by the recent gene duplication pair *achi/vis* are virtually identical to each other and are homologues of the human TGIF sequence-specific DNA-binding factor (Ayyar et al., 2003; Wang and Mann, 2003). Both *achi* and *vis* are expressed in many cells throughout development. However, flies that are null mutant for both genes are viable but male sterile, with a meiotic arrest phenotype. Achi/Vis proteins are localized to chromatin in wild-type primary spermatocytes and in *aly* mutant testes (Wang and Mann, 2003).

Despite lacking any predicted DNA-binding motifs, both Aly and Comr proteins are concentrated on chromatin in primary spermatocytes. However, the nuclear localizations of Aly and Comr are mutually dependent: Aly remains cytoplasmic in *comr* mutant testes and vice versa. By contrast, the localization of Aly and Comr to the nucleus is independent of *achi/vis* (Ayyar et al., 2003; Jiang and White-Cooper, 2003; Wang and Mann, 2003; White-Cooper et al., 2000). Aly, Comr and Achi/Vis co-immunoprecipitate from testis protein extracts, suggesting that they interact in a common complex in vivo (Wang and Mann, 2003).

We describe the identification, cloning and characterization of the *Drosophila* *aly*-class meiotic arrest gene, *matotopetli* (*topi*). The similarity in phenotype between *topi* and *aly*, *comr* and *achi/vis* suggests that these genes act together in a common pathway. *topi* encodes a testis-specific predicted Zn-finger

protein, making *Topi* the second putative sequence-specific transcription factor to be placed into the *aly* class. At least two separate Zn-finger containing regions of *Topi* are sufficient to bind directly to a 100 amino acid region in the middle of the Comr protein. Like *achi/vis*, *topi* activity is not required for the nuclear localization of Aly and Comr proteins. We suggest that the DNA-binding activities of *Topi* and Achi/Vis are required to recruit Aly and Comr to the promoters of target genes in primary spermatocytes. Consistent with this idea, the majority of target genes require both the *topi* and *achi/vis* DNA-binding factors for full transcriptional activation. However, rare exceptions to this rule have been revealed via microarray analysis. A small number of target promoters rely to a much greater extent on one or other of these DNA-binding activities, indicating that in at least some contexts the activities of *achi/vis* and *topi* are independent.

## Materials and methods

### Fly strains and husbandry

Flies were raised on standard cornmeal molasses (or sucrose) agar at 25°C. Visible markers and balancer chromosomes are described in FlyBase except where otherwise noted (FlyBase, 1999). *red e* or *bw*; *st/TM6B* was used as the wild-type strain. *aly*<sup>5</sup> *red e*/ *TM6C*, *w*; *can*<sup>3</sup> *red e*/ *TM6C*, *cn achi*<sup>Z3922</sup>*vis*<sup>Z3922</sup> *bw*/ *CyO*, *cn comr*<sup>Z1340</sup> *bw*/ *CyO*, *mia st*/ *TM6B* and *sa*<sup>1</sup> *red*/ *TM3* stocks were used to generate homozygotes for comparison to other meiotic arrest loci (Ayyar et al., 2003; Jiang and White-Cooper, 2003; Lin et al., 1996). *topi*<sup>Z3-2139</sup>, *topi*<sup>Z3-0707</sup> and *topi*<sup>Z3-3767</sup> were provided by C. Zuker from his collection of EMS-generated, viable lines, screened for male sterility by B. Wakimoto and D. Lindsley.

### Microscopy and immunofluorescence

Live testes were dissected, squashed in 4 µg/ml Hoechst 33342 in testis buffer (183 mM KCl, 47 mM NaCl, 10 mM Tris pH 6.8) and examined by phase contrast and fluorescence microscopy. *topi* mutant testes contained many balls of cells, and no elongating cysts, hence the name *matotopetli*, which means 'balls' in the Aztec language, nahuatl. Aly and Comr proteins were visualized using a BioRad Radiance Plus confocal microscope, using rabbit anti-Aly or rabbit anti-Comr antibodies detected with FITC-conjugated secondary antibodies (Sigma), DNA was co-stained with propidium iodide (Jiang and White-Cooper, 2003; White-Cooper et al., 2000). Phase contrast images were captured using a Photometrics cooled CCD camera connected to a Zeiss Axiophot microscope, or a Q-imaging Retiga 1300 CCD camera linked to an Olympus BX50 microscope using IP Lab Spectrum or Openlab software (Improvision) and then imported into Photoshop. Images of RNA in situ hybridization to testes were captured on colour slide film, this was scanned and imported into Photoshop.

### Northern blotting and in situ hybridization

Northern blotting of poly-A+ RNA from whole males, whole female, agametic males and embryos using *topi* and *rp49* probes was carried out as described previously (Hiller et al., 2001). RNA in situ hybridization was carried out as previously described (White-Cooper et al., 1998). Dig-labelled RNA probes were generated using dig-RNA labelling mix (Roche). Probes for *topi*, *cyclinA*, *cyclinB*, *twine*, *fzo* and *Mst87F* were generated by transcription off a linearized cDNA-clone plasmid. For *TrxT* and *CG8349* a T3 RNA polymerase promoter was included in the 3' RT-PCR primer, and purified PCR product was used as the template for transcription of the labelled RNA probe.

### Microarray analysis and RT-PCR

Full details of the microarray analysis will be presented elsewhere. Testes and seminal vesicles were dissected from 0- to 1-day-old males

homozygous for *red e* (control), *aly*<sup>5</sup>, *can*<sup>3</sup>, *comr*<sup>z1340</sup>, *achi*<sup>Z3922</sup>*vis*<sup>Z3922</sup> and *topi*<sup>Z3-2139</sup> in testis buffer, and placed at  $-80^{\circ}\text{C}$  within 30 minutes of dissection. For each genotype, three independent RNA samples were made from 200 testes. Testes were homogenized in Trizol (Roche), then shipped on dry ice to the BBSRC IGF facility in Glasgow for RNA extraction, labelling and Affymetrix array hybridization (<http://www.mblab.gla.ac.uk/igf/index.html>). Total RNA (6  $\mu\text{g}$ ) per sample was used for probe synthesis. Normalized signal intensities were averaged over the three replicates and compared between genotypes. Genes were selected for analysis from a list of genes expressed in wild-type testes where the mean signal for *topi* was eight times higher or lower than that for *achi/vis*. *P* values were calculated with a one tailed *t*-test on this pair of samples.

For RT-PCR, total RNA was extracted from dissected testes with Trizol reagent. The RNA was resuspended in RNase free water at a concentration of three testes worth/ $\mu\text{l}$ . First-strand cDNA from 4  $\mu\text{l}$  of this sample was generated using oligo-dT primers with the SuperScript II reverse transcriptase system (Invitrogen) following the manufacturer's instructions. As a negative control, wild-type testis RNA was used as the template and reverse transcriptase was omitted from the reaction. cDNA derived from 0.18 testes was used for each RT-PCR reaction and amplified with Taq DNA polymerase (Qiagen) with 24 amplification cycles. Genomic DNA from wild-type flies was used as a positive PCR control.

PCR primers (synthesized by MWG) were designed to amplify 400-1000 bp fragments from the transcript of the genes of interest. The T3 RNA polymerase promoter site, preceded by 6 bp of random sequence, was incorporated into the 3' primer to facilitate production of RNA probes directly from the RT-PCR product. Primer sequences were: CG8349-5' GCTCCTTCAGCGCTACATGC; CG8349-3'T3 GCAACGAATTAACCCTCACTAAAGGGCGCATAGGCACATCG; TrxT-5' TCGGCGAGGGCAGAGCTC; TrxT-3'T3 GCAACGAA-TTAACCCTCACTAAAGGGCATTCTCGTCGTGGGC.

### Cloning of *topi*

*topi* was mapped by recombination between *th* and *cu*, and further localized by deficiency complementation to polytene interval 85E9-13. The *topi* region was defined by the right breakpoints of *Df(3R)GB104*, which complemented *topi*, and *Df(3R)by10*, which failed to complement *topi*. This identified a 60 kb region of genomic DNA containing 23 predicted genes. We used representation of ESTs from different tissues as a crude guide to the expression pattern of these genes. Seventeen were not represented in the EST set derived from testis. Four had ESTs from both testis and other tissues, and two (*CG8484* and *CG8526*) had ESTs from only testis libraries. The premature stop codons in *topi*<sup>Z3-2139</sup> and *topi*<sup>Z3-0707</sup> and the mis-sense mutation in *topi*<sup>Z3-3767</sup> were identified by PCR amplifying the genomic region containing *CG8484* from homozygous males, and sequencing the bulk PCR product. Predicted Zn-finger motifs within Topi were identified by eye and using the InterPro analysis tool (<http://www.ebi.ac.uk/InterProScan/>). *topi* homologues from *Anopheles gambiae* (on AAAB01008944.1) and *Drosophila pseudoobscura* (on Contig815\_Contig5737) were identified using Blast searches of the genome sequence databases.

### Yeast two hybrid interaction screen

A Comr-Gal4-DNA-binding-domain fusion construct was made by subcloning the ORF from a full-length *comr* cDNA clone into the vector pGBKT7 using *NdeI* (at the start codon) and *NotI*. As no testis cDNA libraries suitable for two-hybrid screening were available, we generated and screened a testis cDNA-Gal4-Activation Domain (AD) fusion protein library by in vivo recombination using the Matchmaker library construction and screening kit (Clontech). Total RNA (1  $\mu\text{g}$ ), isolated from wild-type testes with Trizol, was used to synthesize first-strand cDNA using an oligo d(T) primer. Double-strand cDNA was synthesized with SMART III and CDS III anchors. The AD fusion library construction and two-hybrid screen were carried out in one

step by co-transforming the yeast strain AH109 with ds cDNA, pGADT7-Rec and pGBKT7/*comr*. Colonies were picked from SD/-Ade/-His/-Leu/-Trp/X- $\alpha$ -Gal selection plates after 7 days. A screen of  $10^6$  independent co-transformants yielded 39 colonies that grew under selective conditions and were blue in the presence of X- $\alpha$ -Gal. AD/library plasmids were isolated from each positive yeast colony, transformed into *E. coli* and sequenced; one of the positive clones contained the full ORF of CG8484 – *topi*.

### Deletion analysis plasmid construction

Determination of the protein interaction domains necessitated construction of *comr* and *topi* deletion series. PCR products for the deletion derivatives were subcloned into pGBKT7 using *NdeI* and *NotI* sites engineered into the primers. Co-transformation of HA109 cells was with pGADT7-RecTopi (full length Topi) or pGADT7-RecComr (full length Comr) as appropriate. A 622 bp fragment covering five zinc fingers in Kruppel was generated by PCR from wild-type genomic DNA.

### Transfection, expression and immunoprecipitation in mammalian tissue culture cells

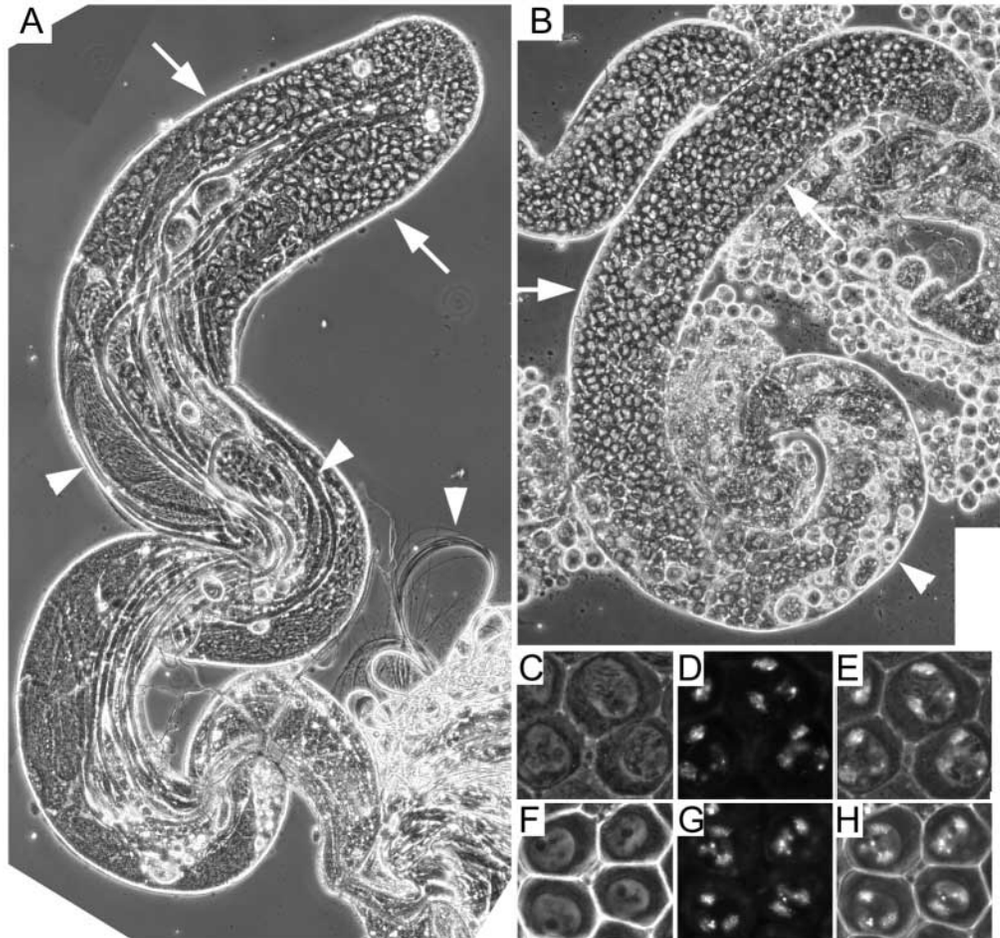
The Topi ORF was subcloned into HA-tagged pCDEF3 and the Comr ORF into FLAG-tagged pCDEF3 respectively at *NdeI* and *NotI* sites. 293T cells were cultured in DMEM with 10% FCS to 90% confluence then co-transfected with 10  $\mu\text{g}$  HA-tagged Topi and Flag-tagged Comr using Lipofectamine 2000 reagent (Invitrogen). As a negative control, 293T cells were co-transfected with HA-tagged Topi and Flag-tagged human Smad1. Forty-eight hours after transfection, cells were collected and processed for immunoprecipitation (Bennett and Alpey, 2002), except that protein G sepharose incubation was carried out at  $4^{\circ}\text{C}$  overnight.

## Results

### Wild type function of *matotopetli* is required for male meiotic division and spermatid differentiation

Mutations in *matotopetli* (*topi*) block both progression of the meiotic cell cycle in males and onset of spermatid differentiation. Wild-type adult *Drosophila* testes contain a spatially organized temporal array of differentiating male germ cells (Fig. 1A) (reviewed by Fuller, 1993). A small population of germline stem cells resides at the testis apical tip. Division of a germline stem cell and associated somatic cyst progenitor cells gives one spermatogonial daughter, which is committed to differentiation and encapsulated by two somatically derived cyst cells. Four mitotic amplification divisions of the spermatogonium results in a cyst of 16 interconnected primary spermatocytes, which then enter an extended period of growth and gene expression followed by meiotic division to produce a cyst of 64 round spermatids. By a programme including extensive elongation and other morphological changes, the spermatids differentiate into mature sperm. Testes lacking *topi* function contained germ cell cysts up to and including mature primary spermatocytes, but lacked cells in meiotic division or spermatid differentiation stages, placing *topi* into the meiotic arrest class of genes (Fig. 1B). *topi* null flies were viable and female fertile. Three alleles of *topi* were identified by screening a collection of EMS induced, viable, male sterile mutations (generated by E. Koundakjian and C. Zuker, screened for male fertility by B. Wakimoto and D. Lindsley). *topi*<sup>Z3-2139</sup> was identical in phenotype to *topi*<sup>Z3-0707</sup>. *topi*<sup>Z3-3767</sup> appeared to be a hypomorphic allele, as occasional progression of mature primary spermatocytes into the meiotic divisions was observed in homozygotes and hemizygotes (data not shown).

**Fig. 1.** The *topi* meiotic arrest phenotype. (A) Phase-contrast image of wild-type whole testis, apical tip in top right-hand corner. Primary spermatocytes occupy most of the apical end (arrows). Elongating spermatid bundles (arrowheads) are seen inside the testis, and spilling out from the distal end. (B) Phase-contrast image of *topi*<sup>Z3-2139</sup> whole testis. Only stages up to mature primary spermatocytes (arrows) are seen. Degenerating primary spermatocytes are present in the distal region of the testis (arrowheads). (C-H) High-magnification views of mature spermatocytes from squashed preparations of wild-type (C-E) and *topi*<sup>Z3-2139</sup> (F-H) testes viewed by phase-contrast (C,F) and Hoechst DNA staining (D,G) or both visualization methods simultaneously (E,H).



The mature spermatocytes present in *topi*<sup>Z3-3767/Df(3R)by10</sup> or *topi*<sup>Z3-3767</sup> homozygous mutant testes resembled wild-type mature primary spermatocytes by phase contrast microscopy (Fig. 1C,F). However the chromatin morphology of mutant spermatocytes was subtly different from wild type when visualized by inclusion of Hoechst 33342 in the squash buffer. Mature wild-type primary spermatocytes have decondensed chromosomes, which are typically found as three chromatin domains close to the nuclear membrane (Fig. 1D,E). As primary spermatocytes pass through the G2-Meiosis I transition, the chromosomes condense and move away from the nuclear envelope. Chromosomes in *topi* arrested cells were somewhat more compact in appearance than in mature wild-type cells in late G2 (Fig. 1G,H), suggesting a block in the G2-M transition of meiosis I in *topi* mutant males. *topi* mutant chromatin most closely resembled that found in *achi/vis* mutant testes, and did not show the fuzzy, decondensed morphology typical of *aly* and *comr* (Ayyar et al., 2003; Jiang and White-Cooper, 2003).

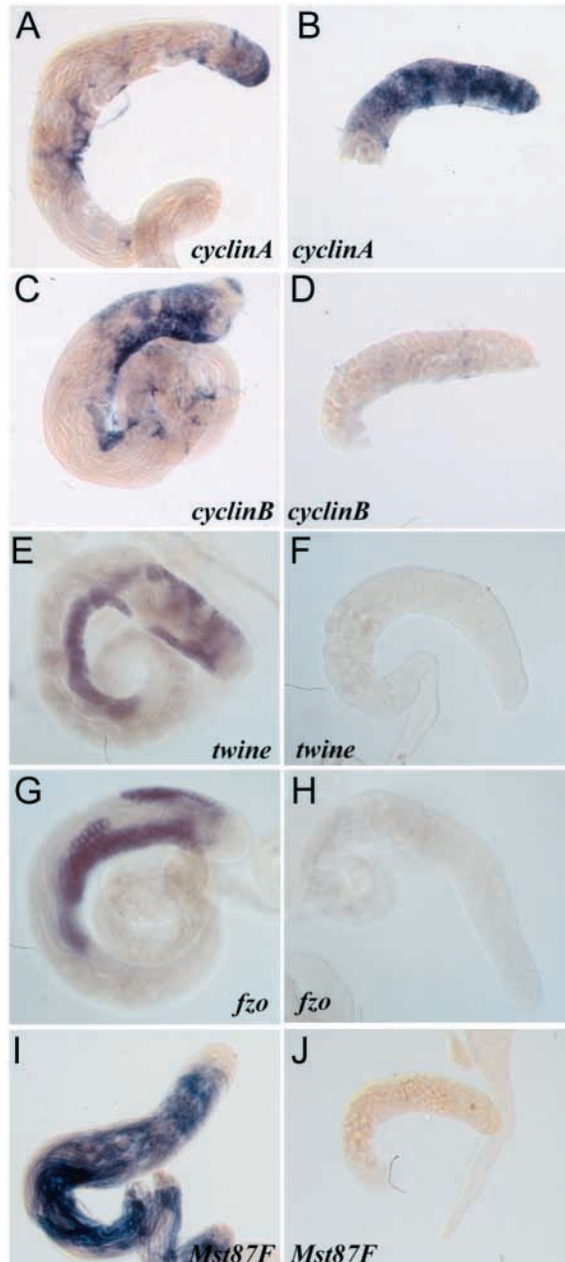
#### ***matotopetli* belongs to the *aly*-class of meiotic arrest genes**

RNA in situ hybridization analysis of the effects of *topi* mutants on transcription of meiotic cell cycle and spermatid differentiation genes placed *topi* into the *aly* class of meiotic arrest mutants. The two strong alleles of *topi* showed greatly

reduced levels of expression in primary spermatocytes of the meiotic cell cycle transcripts *twine*, *cyclin B* (Fig. 2C-F) and *boule* (not shown). The transcript levels of several spermatid differentiation genes, including *fzo* and *Mst87F*, were also dramatically reduced compared with wild-type levels (Fig. 2G-J). Some expression of *fzo* was detected in *topi*<sup>Z3-3767</sup> mutant testes, consistent with the weaker phenotype seen in this allele by phase-contrast microscopy (data not shown).

#### ***matotopetli* function is not required for normal association of Aly and Comr proteins with chromatin in primary spermatocytes**

Our phenotypic analysis indicated that *topi* is a new member of the *aly*-class of meiotic arrest genes, which includes *aly*, *comr* and *achi/vis*. All three of these proteins localize to nuclei of wild-type primary spermatocytes. The behaviour of Aly and Comr proteins in *achi/vis* mutant testes and *topi* mutant testes was extremely similar. Immunofluorescence analysis with anti-Aly and anti-Comr antibodies revealed that both Aly and Comr proteins localized to nuclei in spermatocytes from males carrying strong loss of function mutations in *topi* (Fig. 3). In wild-type primary spermatocytes, Aly and Comr proteins were concentrated on chromatin as well as being distributed throughout the nucleoplasm. By contrast, in *topi* mutant testes, Aly and Comr proteins did not appear to concentrate on chromatin, but were uniformly distributed within the nucleus.



**Fig. 2.** In situ hybridization to whole testes. In situ hybridization to wild-type (A,C,E,G,I), *topi*<sup>Z3-2139</sup> (B,D,J) or *topi*<sup>Z3-0707/Df(3R)by10</sup> (F,H) mutant testes revealed that *topi* is an *aly*-class meiotic arrest gene. The distal ends of some mutant testes were broken off during the staining. Probes were anti-sense transcripts from *cyclinA* (A,B), *cyclinB* (C,D), *twine* (E,F), *fzo* (G,H) or *Mst87F* (I,J).

***matotopetli* encodes a Zn finger protein expressed specifically in primary spermatocytes**

The *topi* gene was localized by a combination of recombination mapping and deficiency complementation (see Materials and methods). Sequence analysis of genomic DNA fragments amplified from *topi*<sup>Z3-3767/Df (3R)by10</sup>, *topi*<sup>Z3-2139</sup> and *topi*<sup>Z3-0707</sup> mutant animals by PCR revealed that each of the three *topi* alleles caused lesions in *CG8484*, identifying this

predicted gene as *matotopetli*. *topi*<sup>Z3-0707</sup> and *topi*<sup>Z3-2139</sup> introduced stop codons, in the first third and in the last third of the predicted protein, respectively. The weaker allele (*topi*<sup>Z3-3767</sup>) carried a mis-sense mutation.

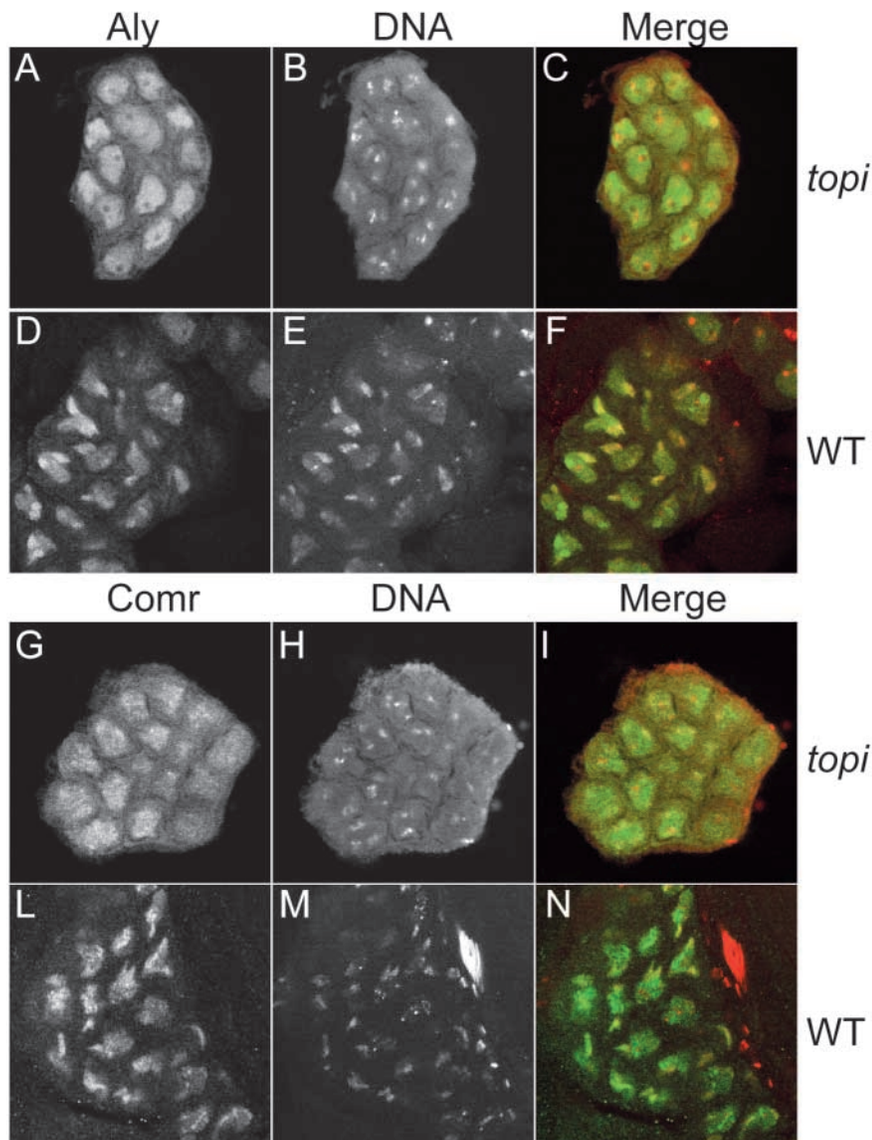
*topi* encodes a 814 amino acid multiple Zn-finger predicted protein with a predicted molecular weight of 92 kDa (Fig. 4). A near full length cDNA isolated from testes (AT25463) was identified in the BDGP EST project and was fully sequenced. The conceptually translated Topi protein contains 10 C2H2 and one C2HC class Zn fingers predicted by the PFAM and prosite protein domain analysis tools. The predicted Zn fingers are clustered in the central region of the protein (amino acids 230-650), with the first two fingers being separated from the remaining nine by a 64 amino acid spacer. The mis-sense mutation in *topi*<sup>Z3-3767</sup> alters a conserved residue, G(516)D, just after the second Zn-binding cystine in the seventh predicted Zn finger. The non-sense mutation in *topi*<sup>Z3-0707</sup> would truncate the protein after the first predicted Zn finger (at amino acid 261). The non-sense mutation in *topi*<sup>Z3-2139</sup> would truncate the predicted Topi protein within the seventh predicted Zn finger (amino acid 521), potentially encoding a protein with six Zn fingers. Blast searches were used to identify putative *topi* orthologues, with sequence conservation extending beyond the Zn finger domains, in the mosquito *Anopheles gambiae* and in *Drosophila pseudoobscura*. Unlike both fly genes, the mosquito protein had only 10 predicted Zn fingers owing to lack of three crucial Zn-binding residues from the second Zn finger.

The *matotopetli* gene encodes a 2.7 kb transcript that was detected in northern blots of polyA+ RNA from whole adult males but not from adult males lacking a germ line, females or embryos (Fig. 5A). RNA in situ hybridization to whole testes revealed that *topi* transcript was expressed specifically in primary spermatocytes. *topi* transcript was not detected in spermatogonia or elongating spermatids (Fig. 5B). *topi* transcript was low or not detectable in spermatocytes from *topi*<sup>Z3-0707/Df</sup> males (Fig. 5C).

***matotopetli* protein physically interacts with Comr**

A full-length clone of the *topi* predicted Zn-finger protein was identified independently in a yeast-two hybrid screen for testis cDNAs encoding proteins that interact with Comr (see Materials and methods). Full-length Comr did not interact in the two-hybrid system with a domain containing five C2H2 Zn fingers from the transcriptional repressor protein Kruppel, indicating that the Comr-Topi interaction is specific. The interaction between Topi and Comr proteins was confirmed using co-expression and immunoprecipitation from mammalian tissue culture cells (Fig. 6C). FLAG-tagged Comr and HA-tagged Topi were co-expressed in 293T tissue culture cells. HA-tagged Topi protein had an apparent *M<sub>r</sub>* of 92 kDa, correlating well with the predicted protein size. Immunoprecipitation with anti-FLAG antibodies, followed by western blotting with anti-HA antibodies showed that Topi co-immunoprecipitated with Comr. The reciprocal experiment, immunoprecipitation with anti-HA antibodies and blotting with anti-FLAG, also showed co-immunoprecipitation of Comr and Topi. To control for non-specific interactions in the cultured cells we co-expressed HA-Topi with FLAG-Smad1. We were not able to co-immunoprecipitate this pair of proteins.

The domains of Comr and Topi responsible for the



**Fig. 3.** Antibody stain of Aly and Comr in wild type versus *topi*. Immunofluorescence staining of Aly (A,D, green in merge) and DNA (B,E, red in merge) on wild-type (D-F) and *topi*<sup>Z3-2139</sup> (A-C) testes revealed that Aly protein is nuclear in both wild type and mutant, but that Aly fails to accumulate on the chromatin in *topi*<sup>Z3-2139</sup> mutant spermatocytes. Immunofluorescence staining of Comr (G,L, green in merge) and DNA (H,M, red in merge) on wild-type (L-N) and *topi*<sup>Z3-2139</sup> (G-I) testes similarly revealed that Comr protein is nuclear in both wild type and mutant, but that Comr fails to accumulate on the chromatin in *topi*<sup>Z3-2139</sup> mutant spermatocytes.

*topi* and *achi/vis*. However, transcription of a subset (about 42) of genes was much more dependent on *achi/vis* than on any of the other meiotic arrest loci. Transcription of a different subset (about 44) of genes seemed to be much more dependent on *topi*. The raw and log ratio array data for these genes are presented in Table S1 at <http://dev.biologists.org/supplemental/>, along with data for genes whose expression in wild-type and meiotic arrest mutant testes has already been examined by RNA in situ hybridization or northern blotting (White-Cooper et al., 1998). Further analysis of representative genes by RT-PCR from wild-type and mutant testes confirmed the *achi/vis* versus *topi* differential dependence.

*CG8349* encodes a predicted cytidine deaminase. 11 testis ESTs for this gene have been sequenced, no *CG8349* ESTs have been sequenced from other tissue libraries, indicating that *CG8349* transcription is highly testis enriched. Microarray analysis indicated that expression of this gene was approximately five times lower in *aly*, *comr*

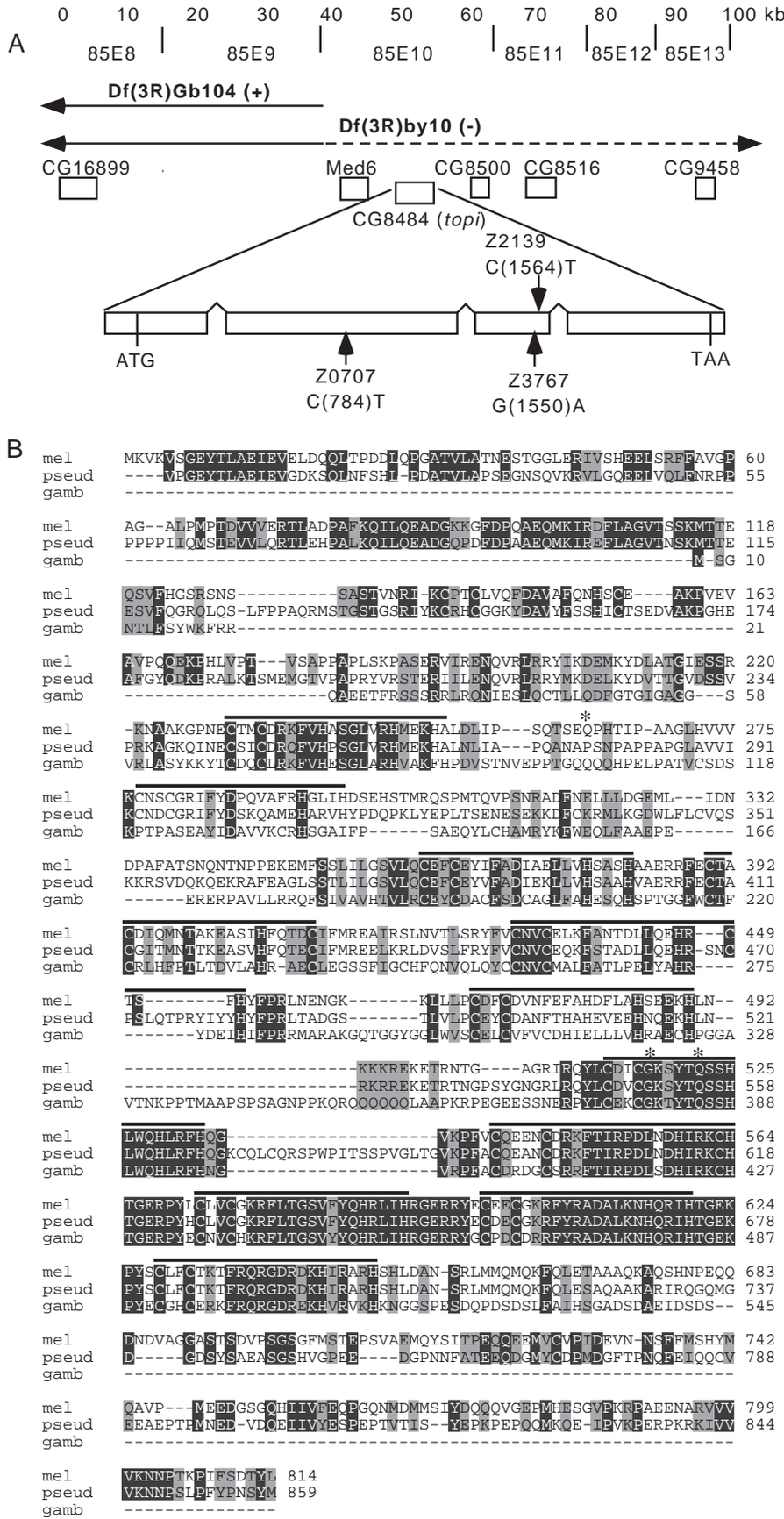
and *topi* than in wild type, while expression of *CG8349* was reduced by two-hundred times in *achi/vis* compared with wild type. By RT-PCR, we found that *CG8349* transcript was detectable, but levels were dramatically reduced in *aly*, *comr*, *topi*, *mia* and *sa* mutant testes. However, no transcript was detected in *achi/vis* mutant testes (Fig. 7A). To further investigate these changes in gene expression, we used in situ hybridization against wild-type and mutant testes. The relative level of transcript can be compared between wild-type and mutant testes using this method as the tissues were mixed before hybridization. In situ hybridization revealed that *CG8349* was expressed in primary spermatocytes, and the transcript persisted into mid-late stages of spermatid elongation (Fig. 7B). As predicted by the RT-PCR and array analysis, the in situ hybridization signal intensity for *CG8349* was lower than wild type in *aly*, *comr* and *topi* testes (Fig. 7C,E), and the transcript was undetectable by in situ hybridization in *achi/vis* mutant testes (Fig. 7D).

#### Differential requirements for *topi* and *achi/vis* for expression of some target genes

To explore whether *topi* and *achi/vis* might differ in their target gene specificities, we carried out a set of DNA microarray experiments to examine gene expression in wild-type and meiotic arrest mutant testes (see Materials and methods) (H.W.-C., unpublished). About 1000 genes were expressed at reduced levels (at least fourfold) in *aly*-class meiotic arrest mutants than in wild type, most of these transcriptional targets of the meiotic arrest genes depended equally on the functions of *aly*, *comr*,

and *topi* than in wild type, while expression of *CG8349* was reduced by two-hundred times in *achi/vis* compared with wild type. By RT-PCR, we found that *CG8349* transcript was detectable, but levels were dramatically reduced in *aly*, *comr*, *topi*, *mia* and *sa* mutant testes. However, no transcript was detected in *achi/vis* mutant testes (Fig. 7A). To further investigate these changes in gene expression, we used in situ hybridization against wild-type and mutant testes. The relative level of transcript can be compared between wild-type and mutant testes using this method as the tissues were mixed before hybridization. In situ hybridization revealed that *CG8349* was expressed in primary spermatocytes, and the transcript persisted into mid-late stages of spermatid elongation (Fig. 7B). As predicted by the RT-PCR and array analysis, the in situ hybridization signal intensity for *CG8349* was lower than wild type in *aly*, *comr* and *topi* testes (Fig. 7C,E), and the transcript was undetectable by in situ hybridization in *achi/vis* mutant testes (Fig. 7D).

The microarray analysis indicated that transcripts of *TrxT*, a testis specific isoform of thioredoxin (Svensson et al., 2003),



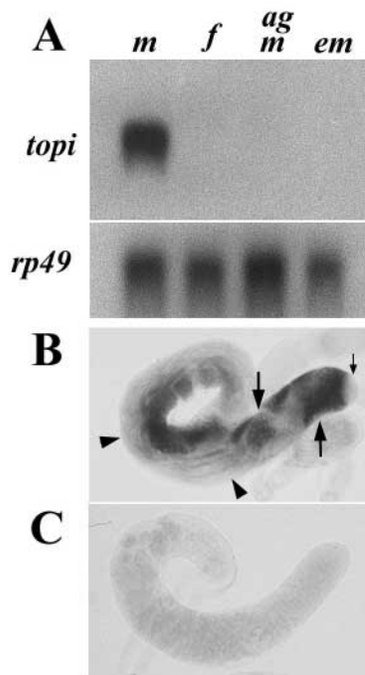
**Fig. 4.** Molecular cloning of *topi*. (A) *topi* genetic region was defined by the right breakpoints of Df(3R)Gb104 and Df(3R)by10, an interval of about 60 kb, which contains 23 predicted genes of which five are shown. *topi* gene structure and the location of lesions in mutant alleles are shown. (B) *topi* predicted amino acid sequence (mel) aligned with the putative *topi* orthologues from *D. pseudoobscura* (pseud) and *A. gambiae* (gamb). Sequence identity between all three proteins is indicated by black boxes, sequence similarity is indicated by grey boxes. Predicted Zn-finger motifs are indicated with lines above the sequences. The amino acids affected in the mutant alleles are indicated with an asterisk above the sequence.

were 2.5 times less abundant in *aly* and *comr* than in wild-type or *achi/vis* mutant testes. However, *TrxT* transcript was at least fifty times less abundant in *topi* mutant testes than in wild type. Consistent with these data, we were unable to RT-PCR *TrxT* from *topi* mutant testes, while the RT-PCR of *TrxT* revealed only a mild reduction in transcript levels compared with wild type in *aly*, *comr* and *achi/vis* mutant testes (Fig. 7F). No reduction in transcript level was evident by RT-PCR in *mia* or *sa* mutant testes. *TrxT* expression was first detected by in situ hybridization in primary spermatocytes in wild-type testes, the transcript persisted after the meiotic divisions and was detected until mid-elongation stages of spermiogenesis (Fig. 7G). *aly* and *comr* mutant testes showed expression in the primary spermatocytes; however, the signal intensity in these cells was lower than in the wild-type control (Fig. 7H). Thus, although significant expression of *TrxT* was detected in *aly* and *comr* testes, *aly* and *comr* are both required for full expression of *TrxT*. Signal intensity was similar to control levels in *achi/vis* mutant testes (Fig. 7I). *TrxT* transcript was undetectable in *topi* mutant testes by in situ hybridization (Fig. 7J), confirming that *topi* is absolutely required for expression of *TrxT*.

## Discussion

### A pathway for recruitment of the *aly*-class meiotic arrest proteins to chromatin

The *aly* class meiotic arrest genes of *Drosophila* form a functional module required for transcription in primary spermatocytes of a number of meiotic cell



**Fig. 5.** *topi* is expressed specifically in primary spermatocytes. (A) Northern blot with *topi* (top) and *rp49* (bottom) probes. A 2.7 kb *topi* transcript was detected in RNA from whole males (m), but not from females (f), agametic males (ag m) or embryos (em). *rp49* was detected at similar levels in all the samples. (B) RNA in situ hybridization with *topi* probe to wild-type testis reveals expression in primary spermatocytes (arrows) and absent from spermatogonia (small arrow) and spermatids (arrowheads). (C) RNA in situ hybridization with *topi* probe on *topi*<sup>Z3-0707/Df(3R)by10</sup> mutant testis indicates lack of *topi* transcript.

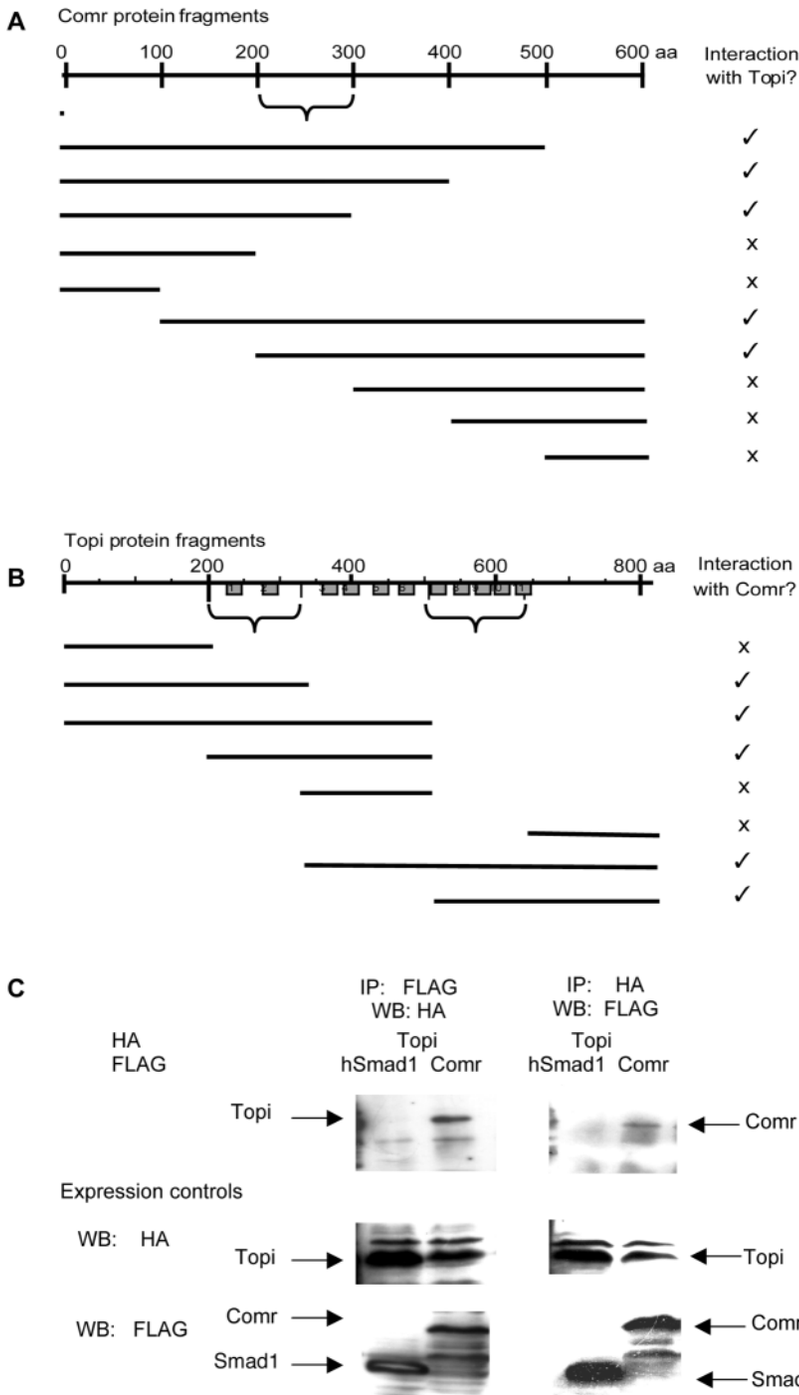
cycle regulatory components and spermatid differentiation genes. *aly* encodes a homologue of the *C. elegans* SynMuvB gene *lin-9*, which is thought to be involved in recruiting the NURD nucleosome remodelling and histone deacetylase complex to repress target genes involved in vulval development (Beitel et al., 2000; White-Cooper et al., 2000). The *Drosophila* Aly protein is found in a complex with Comr, a novel protein that also associates with chromatin in *Drosophila* primary spermatocytes (Jiang and White-Cooper, 2003; Wang and Mann, 2003) (J.J. and H.W.-C., unpublished). A pathway by which Aly and Comr are assembled onto chromatin in primary spermatocytes is now beginning to emerge (Fig. 8). Interaction between Aly and Comr is required for each of the two proteins to localize to the nucleus, because absence of Aly leads to cytoplasmic accumulation of Comr and vice versa (Jiang and White-Cooper, 2003). Once the Aly/Comr complex has entered the nucleus, wild-type function of the predicted DNA-binding proteins Topi and Achi/Vis appears to be required for concentration of the Aly/Comr complex on chromatin. The Aly/Comr complex interacts structurally with Achi/Vis in primary spermatocytes (Wang and Mann, 2003) and Topi directly binds Comr. We propose that the Zn-finger protein Topi, via its physical interaction with Comr, and the TALE-class homeodomain proteins Achi/Vis, via their physical interaction with Aly/Comr, recruit an Aly/Comr complex to chromatin. We propose that a typical target gene,

e.g. *dj*, has both *topi* and *achi/vis* target sequences in its promoter. Topi and Achi/Vis would bind to these regions, either independently or co-operatively, and recruit a pre-formed Aly/Comr complex from the nucleoplasm onto the promoter. Both Topi and Achi/Vis would need to be present to efficiently recruit Aly/Comr. Alternatively, Topi and Achi/Vis could bind to an Aly/Comr complex in the nucleus, with the target sequence for either DNA-binding protein sufficient to localize the entire assemblage to the target promoter. The *aly*-class meiotic arrest gene *topi* plays a crucial role in function of this regulatory module required for activation of a novel transcriptional programme in primary spermatocytes. Loss of function of *topi* results in the arrest of mutant cells as mature primary spermatocytes, and wild-type function of *topi* is required to generate a store of transcripts to support meiosis and the post-meiotic morphological changes that occur during spermatid differentiation.

### Structure of target promoters

Although the meiotic arrest genes are crucial for transcription of a large suite of primary spermatocyte specific genes, they are not global regulators, as many transcripts accumulate to normal levels in the primary spermatocytes of mutant testes. Cis-acting sequences at target promoters would make them dependent on the function of *aly* and *comr*. Targeting of the Aly/Comr complex activity to particular sites on the genome may be crucial for understanding the molecular mechanism of this genetic module. The identification of two potential DNA-binding activities within the *aly*-class of meiotic arrest genes raises questions about the requirements for each of these proteins at different target gene promoters. In our microarray analysis, all genes we found to depend on *aly* or *comr* were also dependent on *topi* and/or *achi/vis*. The overwhelming majority of *topi*-dependent genes are also dependent on *aly*, *comr* and *achi/vis* for their full activation; genes in this class include *Mst87F*, *dj* and *fzo*. This indicates that both Topi and Achi/Vis are usually needed at the promoters of meiotic arrest target genes for full activity. Many transcription factors function as homo- or heterodimers, and many developmentally regulated promoters have binding sites for more than one transcription factor. These characteristics allow a combinatorial mechanism to ensure spatiotemporal accuracy of the transcription of any particular gene. The simplest explanation for the requirement for both Topi and Achi/Vis is that the promoters of the majority of terminal differentiation genes have binding sites for both factors. *achi* and *vis* are expressed at many stages of development, yet their testis target genes remain silent in these other tissues. This silence is likely to be due, at least in part, to the lack of *topi* expression in other tissues. Thus, the cooperation between these two transcription factors is essential for the developmental accuracy of expression of genes required for spermatogenesis.

It is intriguing that transcription of some genes (including *TrxT* and *CG8349*) showed a strong requirement for *topi* but not *achi/vis* (or vice versa), and that expression of these genes was less strongly dependent on the activity of the other meiotic arrest genes. This indicates that, at least in these contexts, the binding of the Achi/Vis and Topi transcription factors to DNA is independent. Another feature these genes have in common is that they are members of clusters of related genes, with different expression patterns. *TrxT* is found adjacent in the



**Fig. 6.** Structural interaction between Topi and Comr proteins. (A) Mapping domains of Comr responsible for the interaction with full-length Topi. Solid bars represent protein fragments tested in the yeast two-hybrid system against Topi. The region responsible for the interaction lies in the 200-300 amino acid fragment (bracket). (B) Mapping domains of Topi responsible for the interaction with full-length Comr. Zn-finger motifs within Topi are indicated with boxes. Fragments containing either the first two, or the final five, Zn fingers can bind Comr (brackets). (C) Co-immunoprecipitation of Comr and Topi from tissue culture. Cells expressed HA-tagged Topi and FLAG-tagged Comr or Smad1. Binding was assessed by IP-ing with anti-FLAG, and blotting with anti-HA (first pair of lanes) or vice versa (second pair of lanes). Topi and Comr co-immunoprecipitate in both cases (top). Total protein expression was assessed by western blotting of the cell lysate (bottom).

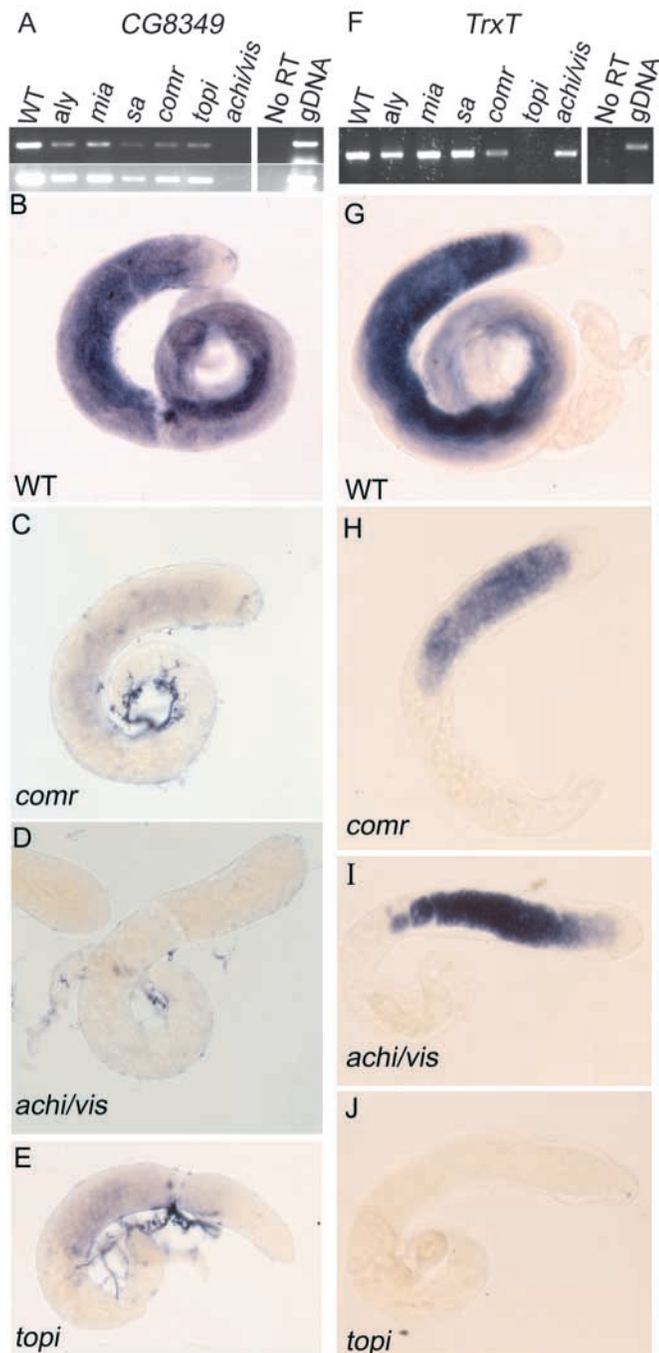
**Functional domains within the Topi protein**

The meiotic arrest gene *topi* encodes a protein with multiple Zn fingers. The final five Zn-fingers of Topi form an evolutionarily constrained unit, which we suggest may function via sequence specific DNA binding activity. *topi* has a large number of transcriptional targets (H.W.-C., unpublished). If, as is likely, Topi binds directly to *cis* acting regulatory elements associated with target genes in a sequence-specific manner, we would expect the protein sequence in the DNA-binding domain to be evolutionarily constrained. Changes in the affinity of Topi for any particular DNA sequence would have to be compensated for by changes in the sequence of a great many promoters, and are therefore likely to be selected against. Conservation of *topi* protein sequences within Diptera is extremely high across the final five Zn-finger motifs, and much lower across the first six Zn-fingers and at the termini. This conservation is particularly apparent in the loop regions of the final five Zn fingers, which are likely to be exposed, and important for interaction with DNA. Additionally these final five Zn finger motifs are more similar to those found in many transcription factors from other taxa. *topi*<sup>Z3-2139</sup> has a non-sense mutation that would truncate the protein before this predicted DNA binding

genome to the female specific thioredoxin, *deadhead* (Svensson et al., 2003). *CG8349* is the 3'-most of a cluster of three predicted cytidine deaminase genes (the others being *CG8360* and *CG8353*). The structure of these promoters may differ from the canonical structure for meiotic arrest target genes, perhaps by having several binding sites for only one of the *aly*-class transcription factors. In this situation partial activation of gene expression would be achieved on binding of one transcription factor. Further recruitment of Aly and Comr (and usually the other DNA-binding factor) would be needed for the robust expression levels seen in wild type.

region, and would presumably therefore be unable to bind DNA. The mis-sense mutation in the hypomorphic allele *topi*<sup>Z3-3767</sup> may reduce the affinity of the protein for its target site, but not totally abolish DNA-binding activity, thus allowing some residual function.

We propose that the first Zn finger of Topi is most important for the binding to Comr. Zn-finger motifs were first identified as nucleic acid binding domains. However Zn fingers are also known to be important for some protein-protein interactions (Evans and Hollenberg, 1988; Tsai and Reed, 1998). There is precedent among known Zn-finger transcription factors, as we



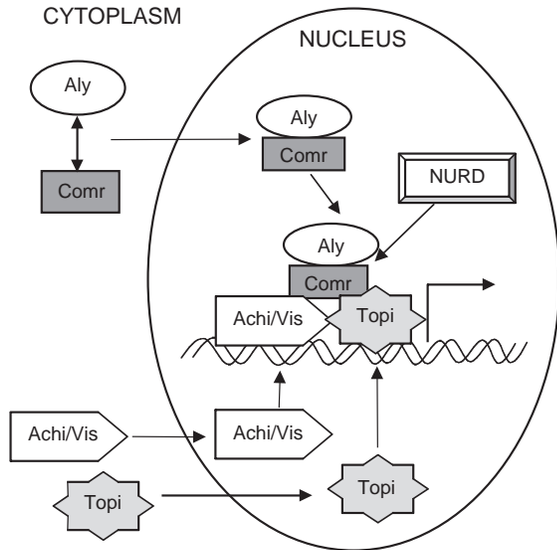
**Fig. 7.** Differential requirement for *topi* and *achi/vis* in expression of some target genes. (A) RT-PCR of *CG8349* from wild-type and meiotic arrest mutant testes. No-RT is a negative control, gDNA is PCR on genomic DNA from wild-type flies. Two exposures of the same gel are shown to emphasize the differences between the mutants and wild type (top), and the lack of product in *achi/vis* (bottom). (B-E) In situ hybridization of *CG8349* probe to testes revealed strong expression in wild-type (B) primary spermatocytes, persisting until mid-late elongation stages, weaker expression in *comr* (C) and *topi* (E), and no detectable signal in *achi/vis* (D) testes. (F) RT-PCR of *TrxT* from wild-type and meiotic arrest mutant testes. No-RT is a negative control, gDNA is PCR on genomic DNA from wild-type flies. (G-J) In situ hybridization of *TrxT* probe testes revealed strong expression in wild-type (G) primary spermatocytes and early-mid elongation stages. The signal in *achi/vis* testes was of a similar intensity to wild type (I). Weaker expression was seen in *comr* (H) and there was no detectable signal in *topi* (J) testes.

binding activity in the yeast two-hybrid system. This interaction may stabilize, or increase the binding affinity of, the Topi-Comr complex. Strikingly, the N-terminal Comr binding region of *topi* is much less well conserved than the last five Zn finger containing domain, even among Diptera (*D. melanogaster*, *D. pseudoobscura*, *A. gambiae*). Also apparent is the relatively low conservation within the loop regions of the Zn fingers. If *comr* is rapidly evolving, the Comr interaction domains within Comr-binding proteins would also be expected to show low levels of conservation. It is possible that the N-terminal Zn-finger Comr-binding domain of Topi has evolved rapidly in concert with the rapid evolution of Comr. Searches of sequence databases had previously indicated failed to identify a homologue of *D. melanogaster comr* in the genome sequence of *A. gambiae*, which diverged from *Drosophila* 250 Mya (Zdobnov et al., 2002). Partial sequence of a *comr* homologue (H.W.-C., unpublished) was found in the set of unassembled genome sequence contigs from another Drosophilid, *D. pseudoobscura*, which diverged from *D. melanogaster* 46 Mya (Bergman et al., 2002). The conservation with *D. melanogaster comr* (31% identical, 52% similar) was remarkably low considering the relatedness of the species. This relative lack of conservation was apparent even in the Topi-binding region.

#### Transcriptional activation by the *aly*-class genes

The exact mechanism by which the *aly* homologue *lin-9* functions is not well understood. However, some insight has been provided by the molecular analysis of other genes in the *C. elegans* Syn-MuvB pathway, including *lin-35* (Rb, Retinoblastoma), *lin-53* (RbAp48) and *hda-1* (histone deacetylase) (Lu and Horvitz, 1998). Other components of the NURD histone deacetylase and chromatin remodelling complex also have SynMuvB activity (Solari and Ahringer, 2000). The biochemical function of NURD is to de-acetylate histones and re-position nucleosomes (Xue et al., 1998; Zhang et al., 1998). Typically, histone de-acetylation is associated with transcriptional repression, although exceptions to this rule have been found (De Rubertis et al., 1996; Struhl, 1998). Another gene in the SynMuvB pathway, *lin-13*, has been shown to encode a putative Rb-interacting protein with multiple C2H2 Zn-fingers, which may be important for targeting Rb and the NURD complex to specific sites

propose for Topi, for certain of the fingers to bind proteins, while others contact the DNA, allowing the transcription factor to target a multi-subunit complex to specific promoters (Kalenik et al., 1997). The Topi protein interacts structurally with Comr, and the region of Topi containing the first two Zn fingers was sufficient for interaction with Comr in yeast two hybrid assays. We propose that the first Zn finger is more important for Comr binding because the second of the two Zn fingers in mosquito Topi lacks crucial Zn-binding residues, so is unlikely to form a conventional finger structure in the folded protein. The final five Zn fingers, which we propose function primarily as DNA-binding motifs, also showed some Comr-



**Fig. 8.** Model for assembly and function of an *aly*-class protein complex at target promoters. Aly and Comr may be imported into the nucleus together or may only form a stable complex in the nucleus. Either protein alone remains in the cytoplasm. Topi and Achi/Vis proteins would enter the nucleus independently. These putative DNA-binding proteins would recognize specific sequences at target promoters, and once bound they would together recruit the Aly/Comr complex to these promoters. Alternatively, Topi and Achi/Vis could assemble with Aly/Comr in the nucleus, and this entire complex would then bind target promoters. This complex would then recruit the NURD histone deacetylase chromatin remodelling complex and promote transcription.

(Melendez and Greenwald, 2000). The loop regions of the Zn fingers in LIN-13 showed no significant similarity with those in Topi, however the two proteins could be playing similar roles in linking *lin-9* and NURD to target promoters. Analogous to the *C. elegans* system, we have proposed that *aly*, and by extension other genes in the same pathway as *aly*, functions through a NURD-complex chromatin-remodelling activity. Many genes require the *aly*-class meiotic arrest genes for activation of expression in primary spermatocytes, while a few may require these genes for transcriptional repression (H.W.-C., unpublished). Several SynMuvB genes have also been shown to be important for expression of transgenes in repetitive extrachromosomal arrays in *C. elegans*, indicating that, at least in certain specific contexts, the SynMuvB pathway activates gene expression (Hsieh et al., 1999). We favour a model in which the putative sequence-specific DNA-binding proteins, Topi and Achi/Vis target Aly and Comr to specific promoters, where Aly/Comr may then recruit a NURD complex to alter local chromatin structure. In the second part of the model, transcription of specific terminal differentiation genes in primary spermatocytes in some way requires the altered chromatin structure set up by the function of the Aly/Comr complex. The *can*-class meiotic arrest genes appear to form a different functional module that is also required for normal levels of transcription of many of the same target genes that require *aly/comr*. So far, all of the *can* class meiotic arrest genes molecularly identified encode homologues of subunits of TFIID. A TFIID complex containing the testis specific TAFs

encoded by the *can*-class meiotic arrest genes may have a higher affinity for the altered chromatin conformation than conventional TFIID, and thus be required for full transcriptional activity.

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