# Requirement of *flex* (*female lethal on X*) in the development of the female germ line of *Drosophila melanogaster*

# ANANYA BHATTACHARYA $^{1\ast},$ S. SUDHA $^1,$ SWATHI BALAKRISHNA $^1$ and H. SHARAT CHANDRA $^{1,2}$

<sup>1</sup>Department of Microbiology and Cell Biology, Indian Institute of Science, Bangalore 560012, India <sup>2</sup>Jawaharlal Nehru Centre for Advanced Scientific Research, Bangalore 560064, India

## Abstract

Drosophila melanogaster females homozygous for flex, an X-linked recessive mutation, do not survive. Hemizygous males are unaffected. Homozygous embryos appear to lack SXL, the product of the Sex-lethal (Sxl) gene, apparently as a result of disruption of Sxl splicing. It is known that both Sxl and its somatic splicing regulators [snf and fl(2)d] also function in the development of the female germ line. For this reason, we investigated the role of flex in the germ line by generating flex/flex clones in flex/+ females. Females carrying such clones in their germ lines do not lay eggs whereas females carrying flex+ eggs lay viable eggs. Additionally, DAPI staining of ovarioles showed that diploid germ cells that are homozygous mutant for flex do not complete oogenesis. These results indicate that the flex<sup>+</sup> gene product may be required for the development of the female germ line.

[Bhattacharya A., Sudha S., Balakrishna S. and Chandra H. S. 1999 Requirement of *flex* (*female lethal on X*) in the development of the female germ line of *Drosophila melanogaster*. J. Genet. **78**, 133–139]

### Introduction

In *Drosophila melanogaster*, the process of sex determination is initiated by a cell-autonomous mechanism which functions transiently in somatic cells of the early embryo (Sanchez and Nöthiger 1983; Cline 1984). This involves measurement of the ratio of the number of X chromosomes (X) to the number of sets of autosomes (A). The X : A ratio activates a special embryonic promoter, *Pe*, of *Sxl* (Keyes *et al.* 1992). At this stage, the initiation of *Sxl* transcription is limited to females; later in embryogenesis, transcription shifts to a maintenance promoter, *Pm*, which is active throughout development in both sexes. SXL, produced from *Sxl*<sub>Pe</sub> mRNA, initiates an autoregulatory feedback loop that directs the productive splicing of transcripts expressed from *Sxl*<sub>Pm</sub> (Bell *et al.* 1991; Bopp *et al.* 1991). *Sxl* controls three subordinate pathways that are necessary for the viability and fertility of the fly: the processes leading to dosage compensation, somatic sexual development and germ line sex determination (for reviews see Baker and Belote 1983; Steinmann-Zwicky *et al.* 1990; Belote 1992; Steinmann-Zwicky 1992a, b; Kuroda *et al.* 1993; Baker *et al.* 1994; Gorman and Baker 1994).

Regulation of sex determination in the germ line is substantially different from that in the soma. Germ line sex determination is regulated by cell-autonomous as well as cell-nonautonomous factors (reviewed by Pauli and Mahowald 1990; Steinmann-Zwicky 1992a, b; Burtis 1993). Female germ cells require the activity of *Sxl* to proceed normally through oogenic differentiation (Schupbach 1985; Nöthiger *et al.* 1989; Steinmann-Zwicky *et al.* 1989). Studies by Bopp *et al.* (1993) have shown that *Sxl* is required for some aspect of germ line development other than establishment and maintenance of sexual identity. The ultimate effect of X : A signals in the germ cells appears to be the same as in the soma: removal of the male-specific exon from *Sxl* pre-mRNA (Bopp *et al.* 1993; Oliver *et al.* 1993). Many of the known components of the X : A signalling elements do

Keywords. sex determination; Sxl; FLP-recombinase; FRT.

<sup>\*</sup> For correspondence. Present address: Waksman Institute, Rutgers University, 190 Frelinghuysen Road, Piscataway, NJ 08854-8020, USA. E-mail: bhatta@waksman.rutgers.edu.

not seem to play any role in the germ line (Schupbach 1985; Granadino et al. 1993; Steinmann-Zwicky 1993). In the sexual differentiation pathway, genes downstream of Sxltra, tra2 and dsx—are not required in female germ cells but rather must function in the surrounding somatic tissues (Marsh and Wieschaus 1978; Schupbach 1982). The tra-dsx signalling pathway may direct differentiation of the female germ line by feminizing the expression of some as yet unknown genes. tra2 is thought to act as a cofactor for some other sex-specific gene which finally activates the somatic signalling pathway (Horabin et al. 1995). The products of  $otu^+$ ,  $ovo^+$ ,  $snf^+$ ,  $fu^+$  and  $fl(2)d^+$  are required for the female-specific splicing of  $Sxl^+$  pre-mRNA and thus for specifying female differentiation of the germ line (Oliver et al. 1988, 1990, 1993; Steinmann-Zwicky 1988; Granadino et al. 1992; Pauli et al. 1993).

During a search for additional X-linked genes affecting sex determination, a female-lethal mutation was isolated in our laboratory from an EMS mutagenesis screen (Anand 1993). Females homozygous for this mutation, named *flex*, do not survive; heterozygous females and hemizygous males are viable and fertile. In contrast to most of the genes affecting sex determination, *flex* does not exhibit female lethality in transheterozygous combination with loss-offunction alleles of *Sxl*, *sisA*, *sisB*, *da*, *snf* and *fl*(2)*d* (A. Anand and H. S. Chandra 1993 Abstract, Proceedings of the XVII International Congress of Genetics, Birmingham, UK).

Three new alleles of *flex* were subsequently isolated:  $flex^2$ , a mutation induced by EMS, and  $flex^3$  and  $flex^4$ , induced by gamma irradiation (Bhattacharya et al. 1999). Investigations on the status of Sxl regulation in flex/flexembryos showed that SXL is not present in such embryos. Also, studies of Sxl transcription suggested that Sxl splicing could be disrupted in homozygous embryos (Bhattacharya et al. 1999). Most splicing regulators of Sxl also function in the female germ line (Oliver et al. 1988; Granadino et al. 1992). It was considered necessary to generate germ line chimaeras to find out whether the  $flex^+$  gene product is essential for the development of the female germ line and whether diploid germ cells homozygous mutant for flex have a recognizable phenotype. Here we report on the generation of flex/flex mitotic clones in flex/+ females with the help of the FLP-FRT recombination system (Chou and Perrimon 1992). Our results showed that the females with *flex/flex* clones lay very few eggs. Also, DAPI staining of dissected ovarioles from these females indicated that oogenesis does not proceed to completion when *flex/flex* clones were generated, suggesting that the  $flex^+$  gene product is apparently required for the development of the female germ line.

Stocks used:  $y w FRT^{9-2}/y w FRT^{9-2}$  (Chou and Perrimon 1992) contains two direct repeats of *FRT* sequences cloned in a P element vector with *white*<sup>+</sup> as selectable marker and *nptII* gene conferring resistance to the antibiotic geneticin. C(1)DX yf  $ovo^{D2}v^{24} FRT^{9-2}/Y$ ; *FLP*<sup>38</sup>/*FLP*<sup>38</sup> (Chou and Perrimon 1992) contains, along with the *FRT* sites, a dominant female-sterile mutation ( $ovo^{D2}$ ). The FLP-recombinase is also cloned in a P element vector under a hsp70 promoter with  $rosy^+$  as a selectable marker. Both these stocks were gifts from Prof. Norbert Perrimon. Among the *flex* alleles, *flex*<sup>2</sup> and *flex*<sup>3</sup> were used in this set of experiments. The description of balancers can be obtained from Flybase.

Generation of germ line clones: The two flex alleles,  $flex^2$  and  $flex^3$ , were recombined onto the chromosome bearing *FRT* at 18E. Females carrying the recombinant chromosome (*flex FRT*) were crossed to males with  $ovo^{D2}$  *FRT*. The progeny of such crosses were given heat shock at appropriate stages of development to induce the synthesis of FLP-recombinase. Recombination occurs between the *FRT* sites leading to the formation of mitotic clones (*flex/flex*) in the developing germ line. The presence of  $ovo^{D2}$  marks the products of recombination. For example,  $ovo^{D2}/+$  is the nonrecombinant genotype, and  $ovo^{D2}/ovo^{D2}$  and *flex/flex* are the recombinant classes. To check the effects of such clones in the female germ line, each of these females was crossed to wild-type males and their egg-laying ability monitored.

DAPI staining of ovaries: (after Warn et al. 1985) The ovaries of each genotypic class of females (3-5 days old) were dissected in cold 1 × EBR (130 mM NaCl, 4.7 mM KCl, 1.9 mM CaCl<sub>2</sub>, 10 mM HEPES, pH 6.9) and transferred to an Eppendorf tube containing  $1 \times EBR$  on ice. EBR was removed and 100 µl of devitellizing buffer [1 vol buffer B (whose composition is given below), 1 vol formaldehyde 36%, 4 vol H<sub>2</sub>O] and 600  $\mu$ l of heptane were added. The mix was shaken vigorously to saturate the buffer with heptane and then gently for 10 minutes at room temperature. This solution was removed and the ovaries were rinsed with  $1 \times PBS$ . The composition of buffer B was 100 mMKH<sub>2</sub>PO<sub>4</sub>/K<sub>2</sub>HPO<sub>4</sub> (pH 6.8), 450 mM KCl, 150 mM NaCl, 20 mM MgCl<sub>2</sub>. Tissues were incubated for 5 minutes at room temperature in  $1 \mu g/ml$  DAPI solution in PBS. They were then rinsed with PBS and mounted in 50 µl glycerol: PBS (50% glycerol). The ovaries were placed on a slide, and individual egg chambers were dissected and examined in the UV channel of a fluorescence microscope.

# Results

#### Generation of flex/flex clones in the germ line

# Flies were maintained on standard cornflour-yeast-sugaragar medium at 25°C unless otherwise stated.

Materials and methods

The effect on oogenesis of a clone of flex/flex cells in the germ line of a flex/+ female was investigated (for details see Materials and methods).  $flex^2$  was recombined onto a

**Table 1.** Clonal analysis of  $flex^2$  in the germ line.

Genotype of females	Stage of heat shock	Per cent egg-laying efficiency*
wf FRT/wf FRT (I)	48–72 hours 72–96 hours	60 (150/250)
w f flex <sup>2</sup> FRT/ovo <sup>D2</sup> FRT (II)	48–72 hours 72–96 hours	12 (8/66) 6 (4/66)
y cho cv v f flex <sup>2</sup> FRT/ovo <sup>D2</sup> FRT (III)	48–72 hours 72–96 hours	3 (2/63) 0 (0/45)
wf FRT/wf FRT & wf flex <sup>2</sup> FRT/ovo <sup>D2</sup> FRT (IV)	Nil	0 (0/85)

\* Expressed as the ratio of the number of females laying viable eggs to the total number of females tested. The detailed description of each of the crosses is given below.

(I) w f FRT/w f FRT females (which served as positive controls) were crossed to  $ovo^{D2} FRT/Y$ ;  $FLP^{38}/FLP^{38}$  males and the progeny were given heat shock at the two stages indicated. The F<sub>1</sub> females were crossed to wild-type males and the egg-laying ability of such females was tested.

(II) w f flex<sup>2</sup> FRT/FM7 females were crossed to  $ovo^{D2} FRT/Y$ ; FLP<sup>38</sup>/FLP<sup>38</sup> males and the progeny were given heat shock as described in the table. Among the F<sub>1</sub> females, the non-Balancer females were collected and their egg-laying ability was monitored as before.

(III) To monitor effect of genetic background  $y cho cvv f flex^2 FRT/FM7$  females were used instead of  $w f flex^2 FRT$ . The crosses are identical to those described in row II.

(IV) As negative controls in this analysis, both  $w f flex^2 FRT/FM7$  females and w f FRT/w f FRT females were used but no heat shock was administered. The crosses were the same as those described earlier.

chromosome carrying the *FRT* insertion at 18E and such recombinant females were crossed to males bearing  $ovo^{D2}$  *FRT* and FLP-recombinase. The progeny were subjected to heat shock at 37°C for two hours at the middle of the second instar (48–72 hours after oviposition) and late third instar (72–96 hours after oviposition). As positive control, w f FRT flies which are flex<sup>+</sup> were also subjected to heat shock to test for the activity of the FLP-recombinase. As negative control, all females (with and without flex<sup>2</sup>) were tested for their egg-laying ability but no heat shock was administered. The results of one such experiment are summarized in table 1.

The frequency of mitotic recombination is around 60% in the positive control ( $flex^+ FRT$ ), which agrees well with published data for  $FLP^{38}$  (Chou and Perrimon 1992). Flies that received no heat shock (i.e.  $flex^+ FRT$  and  $flex^2 FRT$ , which served as negative controls) did not lay any eggs. These data show that there is no leakiness associated with the recombinase and that the clones formed in the heatshocked flies are entirely products of recombination. It was observed that  $flex^2 FRT/ovo^{D2} FRT$  females carrying the flex/flex clones, upon heat shock, lay very few eggs in comparison to  $flex^+ FRT/ovo^{D2} FRT$  females. To examine if there are allele-specific variations in this phenotype, a similar analysis was carried out with  $flex^3$  instead of  $flex^2$ . The egglaying of  $flex^3 FRT$  flies was monitored following heat shock conditions similar to those indicated in table 1. The results are summarized in table 2.

Hence, the observation that females in which presumptive flex/flex clones had been induced lay few or no eggs suggests the possibility that such chimaeric germ lines do not support the formation of viable eggs. To verify the formation of mitotic clones and to see the effect of flex/flex clones on the development of the female germ line, ovarioles from such females were dissected and stained with DAPI.

# Staining of ovaries with DAPI

The ability of germ cells homozygous for *flex* to progress through oogenesis and their capacity to form functional gametes was explored. We had observed that SXL is not present in the somatic tissues of embryos homozygous for *flex* owing to a disruption in the splicing regulation of *Sxl* (Bhattacharya *et al.* 1999). Therefore an attempt was made

**Table 2.** Clonal analysis of  $flex^3$  in the female germ line.

Genotype of females	Stage of heat shock	Per cent egg-laying efficiency*
w f FRT/w f FRT (I)	48–72 hours 72–96 hours	53.2 (83/156)
w f flex <sup>3</sup> FRT/ovo <sup>D2</sup> FRT (II)	48–72 hours 72–96 hours	10.86 (5/46) 8.37 (3/37)
$w f FRT/w f FRT \& w f flex^3 FRT/ovo^{D2} FRT$	Nil	0 (0/40)

\* Expressed as the ratio of the number of females laying viable eggs to the total number of females tested.

Experiments in rows I–III are identical to those described in table 1. Here, instead of  $flex^2$ ,  $flex^3$  have been used to analyse whether any allele-specific variations exist.



**Figure 1.** Dissected egg chambers stained with DAPI: (a) bright field image of a dissected ovary from a wild-type female; (b) typical egg chamber from a wild-type female after staining with DAPI; (c) ovary from an  $ovo^{D2}$ /Bal female showing arrest of oogenesis at stage 6.

to investigate whether flex/flex germ cells mimic the behaviour of diploid germ cells homozygous for loss-offunction mutations of *Sxl*. As in the previous experiment,  $flex^2$  was recombined onto a chromosome carrying the *FRT* insertion at 18E and such recombinant females were crossed to males bearing  $ovo^{D2} FRT$  and FLP-recombinase. The progeny were subjected to heat shock at 37°C for two hours according to the protocol given in the previous experiment. Ovaries from  $flex FRT/ovo^{D2} FRT$  and  $flex^+ FRT/ovo^{D2}$ *FRT* females (with and without heat shock) were dissected and stained with DAPI.

Figure 1a shows the bright field image of a dissected ovary from a wild-type female and 1b a typical egg chamber after staining with DAPI. An ovary from an ovo<sup>D2</sup>/Balancer female showing arrest of oogenesis at stage 6 is shown in figure 1c. It can be seen that  $w f flex^+ FRT/ovo^{D2} FRT$ females which did not receive any heat shock have rudimentary ovaries (figure 2a) compared to ovaries from wildtype females (figure 1). In almost all the ovarioles examined, oogenesis had been arrested at around stage 6. Upon receiving heat shock at 72-96 hours after oviposition, the morphology of the ovarioles changed drastically, and the progress of germ cells following oogenesis could be observed (figure 2b). Oocytes in different stages of development as well as mature eggs could be observed. It will be recalled from the previous experiment that such females are capable of laying viable eggs (tables 1 and 2). The results of DAPI staining of ovaries from these  $flex^+ FRT$  females are thus consistent with the results of the egg-laying experiments.

Similar experiments were carried out with  $flex FRT/ovo^{D2} FRT$  females. The results of staining ovarioles from these females with DAPI are shown in figures 3 and 4. As

described in table 1, both  $w f flex^2 FRT/ovo^{D2} FRT$  and  $v cho cv v f flex^2 FRT/ovo^{D2} FRT$  flies were used to see the effect of genetic background. We observed that flies that received no heat shock had rudimentary ovaries in which oogenesis had been arrested around stage 6 (figures 3a and 4a) as in *flex*<sup>+</sup> *FRT*/*ovo*<sup>D2</sup> *FRT* females without heat shock. Upon heat shock (48-72 hours and 72-96 hours) oogenesis appeared to proceed normally in some ovarioles and immature oocytes could be seen (figures 3b and 4b). But neither mature oocytes nor eggs were observed in any of these ovaries. Similar results were obtained when ovaries from flex<sup>3</sup> FRT/ovo<sup>D2</sup> FRT females were dissected and their nuclei stained with DAPI (data not shown). It will be recalled that in comparison with *flex*<sup>+</sup> *FRT/ovo*<sup>D2</sup> *FRT* females the egglaying ability of these females is drastically reduced (tables 1 and 2). Taken together, these observations indicate that flex/flex clones generated in the germ line of flex/+ females do not give rise to normal eggs. On the other hand, wildtype ( $flex^+$ ) clones generated under identical conditions supported the development of viable eggs. Hence, the  $flex^+$ gene product is probably necessary for the formation of a viable germ line in females.

# Discussion

Germ line chimaeras are invaluable for analysing the tissue specificity (germ line vs somatic) of recessive female-sterile mutations (Wieschaus *et al.* 1981; Perrimon and Gans 1983) as well as for detecting the maternal effect of recessive zygotic lethals (Perrimon *et al.* 1984, 1989). Pole cell transplantation and utilization of a dominant female-sterile



**Figure 2.** Dissected egg chambers stained with DAPI. Ovaries from wf flex<sup>+</sup> FRT/ovo<sup>D2</sup> FRT females were dissected and stained with DAPI: (a) ovary from a female which was not subjected to heat shock; (b) typical egg chamber from a female which had received two hours of heat shock at third instar. Oogenesis proceeded normally in such flies.

mutation (DFS) have previously been employed to address such questions (Illmensee 1973; Perrimon and Gans 1983; Perrimon 1984). A modification of the latter method was made by Chou and Perrimon (1992) in which mitotic exchange was induced with the help of the FLP–FRT system in female germ cells.

*flex* is a zygotic female-lethal mutation (Anand 1993). Experiments aimed at investigating the status of Sxl regulation showed that SXL is absent in *flex/flex* embryos. The absence of SXL is most likely the result of a defect in the female-specific splicing of Sxl (Bhattacharya *et al.* 1999). This raises the possibility that *flex* could be a regulator of Sxl splicing or it could be regulating the function of some other gene which directly affects the splicing of Sxl. Since most of the genes involved in the maintenance of Sxl



**Figure 3.** Dissected egg chambers stained with DAPI. Ovaries from  $w f flex^2 FRT/ovo^{D2} FRT$  females dissected and stained with DAPI: (a) ovary from a female which received no heat shock; (b) a typical egg chamber from a female which, as a third instar larva, had received two hours of heat shock; oogenesis is initiated but neither mature oocytes nor eggs were seen.

activity (snf and fl(2)d) also function in the germ line of the female and they display a characteristic phenotype, it was necessary to find out whether  $flex^+$  gene product is essential for the development of the female germ line, and whether diploid germ cells homozygous for *flex* have a recognizable phenotype.

In the present investigation, the approach of Chou and Perrimon (1992) was used to generate flex/flex germ line clones in flex/+ females. It was observed that females bearing such clones lay few eggs compared to females carrying  $flex^+$  clones generated under similar heat shock conditions (tables 1 and 2). This could be because the  $flex^+$  gene product is necessary for the successful completion of the development of the female germ line. This effect appears to be independent of the genetic background.

To verify the formation of these mitotic clones in the germ line and to see the progress of such germ cells through oogenesis, ovarioles from each of these classes of females (described in table 1) were stained with DAPI. Our experiments showed that flex/flex clones are indeed generated in



**Figure 4.** DAPI staining of dissected egg chambers. Ovaries from females of the genotype  $y cho cvvf flex^2 FRT/ovo^{D2} FRT$  were dissected and stained with DAPI: (a) ovary from a female which did not receive heat shock; (b) typical egg chamber from a female which had received two hours of heat shock as a third instar larva. Here also, oogenesis does not proceed to completion and no mature eggs were seen.

the germ line of the females following heat shock. Although oogenesis appeared to proceed normally in some of these ovarioles, neither mature oocytes nor eggs could be seen. In such ovarioles oogenesis does not proceed beyond stage 10.

We were also interested in the question whether *flex/flex* germ cells mimic the behaviour of diploid germ cells homozygous for loss-of-function mutations of Sxl. Ovaries of females in which the germ cells are  $Sxl^{-}$  are filled with large numbers of small cells that are not morphologically female. They appear either undifferentiated or similar to primary spermatocytes (Schupbach 1985; Oliver et al. 1988, 1990; Steinmann-Zwicky et al. 1989). otu, ovo, fu and snf are required for the female-specific splicing of  $Sxl^+$  premRNA in the female germ line. Diploid germ cells mutant for otu, ovo or snf form tumorous structures which are suppressed by the constitutive expression of  $Sxl^+$  (Steinmann-Zwicky 1988; Oliver et al. 1990; Salz 1992; Pauli et al. 1993). Females transheterozygous for snf and Sxl mutations show sexual transformations, ovarian tumours, and greatly reduced viability. These results led to the proposition that *snf* is needed to activate or maintain *Sxl* expression (Oliver et al. 1988). Additionally, pole cell transplantation experiments have shown that the phenotype of fl(2)d in diploid germ cells resembles that exhibited by  $Sxl^{f4}$  and  $Sxl^{f5}$ , suggesting the likely involvement of fl(2)d in Sxl splicing in the female germ line (Granadino *et al.* 1992).

When ovaries from *flex FRT/ovo<sup>D2</sup> FRT* and *flex<sup>+</sup> FRT/* ovo<sup>D2</sup> FRT females were dissected and their nuclei stained with DAPI, it was observed that the phenotype of the presumptive *flex/flex* clones was different from that exhibited by Sxl/Sxl clones. Our results show that flex affects the development of the female germ line, but *flex/flex* clones neither exhibit an early-arrest phenotype nor form tumorous structures (figures 3b, 4b). In presumptive *flex/flex* clones, the initiation of oogenesis could be observed but neither mature oocytes nor eggs were seen. These results suggest that the requirement of  $flex^+$  is probably not at the stage of establishment of sexual identity in the germ line but at later stages of differentiation. As mentioned before, *flex* appears to be involved in the splicing of *Sxl* in the somatic tissues (Bhattacharya et al. 1999). Thus flex may have different roles in the somatic and germ line tissues. Since we do not yet have a female-viable but sterile allele of *flex*, experiments aimed at deciphering the epistatic relationship between *flex* and other genes involved in germ line sex determination (ovo, otu, snf and others) could not be done. We cannot comment at this stage whether the effect of *flex* on the development of the female germ line is autonomous, since these clones were generated in females which were also heterozygous for ovo<sup>D2</sup>. Hence, it is necessary to generate *flex* homozygous clones in females which do not have any other dominant female-sterile mutation to answer the question whether *flex* acts autonomously in the germ line.

#### Acknowledgements

We thank Prof. Norbert Perrimon and the Bloomington stock centre for several of the fly stocks used in this study, and Dr Anuranjan Anand for providing the  $flex^1$  allele. We also thank Prof. K. VijayRaghavan for discussions and for critical review of the manuscript. We appreciate the assistance given by K. Kuppuswamy in preparing and supplying fly food. This work was supported by grants from the Department of Biotechnology, Government of India.

# References

- Anand A. 1993 A search for additional genes affecting sex determination in *Drosophila melanogaster*. Ph.D. thesis, Department of Microbiology and Cell Biology, Indian Institute of Science, Bangalore, India.
- Baker B. S. and Belote J. M. 1983 Sex determination and dosage compensation in *Drosophila melanogaster*. Annu. Rev. Genet. 17, 345–393.
- Baker B. S., Gorman M. and Marin I. 1994 Dosage compensation in *Drosophila. Annu. Rev. Genet.* 28, 491–521.
- Bell L. R., Horabin J. I., Schedl P. and Cline T. W. 1991 Positive autoregulation of *Sex-lethal* by alternative splicing maintains the female determined state in *Drosophila*. *Cell* 65, 229–239.

- Belote J. M. 1992 Sex determination in *Drosophila melanogaster*: from the X:A ratio to doublesex. *Semin. Dev. Biol.* **3**, 319–330.
- Bhattacharya A., Sudha S., Chandra H. S. and Steward R. 1999 *flex*, an X-linked female lethal mutation in *Drosophila melanogaster*, controls the expression of *Sex-lethal. Development* **126**, 5485–5493.
- Bopp D., Bell L. R., Cline T. W. and Schedl P. 1991 Developmental distribution of female-pecific Sxl proteins in Drosophila melanogaster. Genes Dev. 5, 403–415.
- Bopp D., Horabin J. I., Lersch R. A., Cline T. W. and Schedl P. 1993 Expression of the Sex-lethal gene is controlled at multiple levels during Drosophila oogenesis. Development 118, 797–812.
- Burtis K. C. 1993 The regulation of sex determination and sexually dimorphic differentiation in *Drosophila*. *Curr. Opin. Cell Biol.* 5, 1006–1014.
- Chou T. B. and Perrimon N. 1992 Use of a yeast site-specific recombinase to produce female germ line chimeras in *Drosophila*. *Genetics* **131**, 643–653.
- Cline T. W. 1984 Autoregulatory functioning of a *Drosophila* gene product that establishes and maintains the sexually determined state. *Genetics* **107**, 231–277.
- Gorman M. and Baker B. S. 1994 How flies make one equal two: Dosage compensation in *Drosophila*. *Trends Genet*. **10**, 376–380.
- Granadino B., Juan A. B. S., Santamaria P. and Sanchez L. 1992 Evidence of a dual function in fl(2)d, a gene needed for *Sexlethal* expression in *Drosophila melanogaster*. *Genetics* **130**, 597–612.
- Granadino B., Santamaria P. and Sanchez L. 1993 Sex determination in the germ line of *Drosophila melanogaster*: activation of the gene *Sex-lethal*. *Development* **118**, 813–816.
- Horabin J. I., Bopp D., Waterbury J. and Schedl P. 1995 Selection and maintenance of sexual identity in the *Drosophila* germ line. *Genetics* 141, 1521–1535.
- Illmensee K. 1973 The potentialities of transplanted early gastrula nuclei of *Drosophila melanogaster*. Wilhelm Roux' Arch. Entwickslungmech. Org. **171**, 331–343.
- Keyes L. N., Cline T. W. and Schedl P. 1992 The primary sex determination signal of *Drosophila* acts at the level of transcription. *Cell* 68, 933–943.
- Kuroda M. I., Palmer M. J. and Lucchesi J. C. 1993 X chromosome dosage compensation in *Drosophila*. Semin. Dev. Biol. 4, 107–116.
- Marsh J. L. and Wieschaus E. 1978 Is sex determination in the germ line and soma controlled by separate genetic mechanisms? *Nature* **272**, 249–251.
- Nöthiger R., Jonglez M., Leuthold M., Gerschwiler P. M. and Weber T. 1989 Sex determination in the germ line of *Drosophila* depends on genetic signals and inductive somatic factors. *Development* **107**, 505–518.
- Oliver B., Perrimon N. and Mahowald A. P. 1988 Genetic evidence that *sans fille* locus is involved in *Drosophila* sex determination. *Genetics* **120**, 159–171.
- Oliver B., Pauli D. and Mahowald A. P. 1990 Genetic evidence that *ovo* locus is involved in *Drosophila* germ line sex determination. *Genetics* **125**, 535–550.

- Oliver B., Kim Y.-J. and Baker B. S. 1993 *Sex-lethal*, master and slave: a hierarchy of germ line sex determination in *Drosophila*. *Development* **119**, 897–908.
- Pauli D. and Mahowald A. P. 1990 Germ line sex determination in Drosophila. Trends Genet. 6, 259–264.
- Pauli D., Oliver B. and Mahowald A. P. 1993 The role of the ovarian tumor locus in *Drosophila melanogaster* germ line sex determination. *Development* 119, 123–134.
- Perrimon N. 1984 Clonal analysis of dominant female sterile germ line dependent mutations in *Drosophila melanogaster*. *Genetics* **108**, 927–939.
- Perrimon N. and Gans M. 1983 Clonal analysis of the tissue specificity of recessive female sterile mutations of *Drosophila melanogaster* using a dominant female sterile mutation *Fs*(1)*K*1237. Dev. Biol. 100, 365–373.
- Perrimon N., Engstrom L. and Mahowald A. P. 1984 The effects of zygotic lethal mutations on female germ line functions in *Drosophila. Dev. Biol.* 105, 404–414.
- Perrimon N., Engstrom L. and Mahowald A. P. 1989 Zygotic lethals with specific maternal effect phenotypes in *Drosophila melano*gaster. I. Loci on the X chromosome. *Genetics* **121**, 333–352.
- Salz H. K. 1992 The genetic analysis of *snf*: a *Drosophila* gene required for activation of *Sex-lethal* in both the germ line and the soma. *Genetics* **130**, 547–554.
- Sanchez L. and Nöthiger R. 1983 Sex determination and dosage compensation in *Drosophila melanogaster*: production of male clones in XX females. *EMBO J.* 2, 485–491.
- Schupbach T. 1982 Autosomal mutations that interfere with sex determination in somatic cells of *Drosophila* have no direct effect on the germ line. *Dev. Biol.* **89**, 117–127.
- Schupbach T. 1985 Normal female germ cell differentiation requires the female X chromosome to autosome ratio and expression of *Sex-lethal* in *Drosophila melanogaster. Genetics* **109**, 529–548.
- Steinmann-Zwicky M. 1988 Sex determination in *Drosophila*: the X-chromosomal gene *liz* is required for *Sxl* activity. *EMBO J.* 7, 513–518.
- Steinmann-Zwicky M. 1992a How do germ cells choose their fate, Drosophila as a paradigm. Bioessays 14, 513–518.
- Steinmann-Zwicky M. 1992b Sex determination in *Drosophila* germ cells. *Semin. Dev. Biol.* **3**, 341–348.
- Steinmann-Zwicky M. 1993 Sex determination in *Drosophila: sis-b*, a major numerator element of the X : A ratio in the soma, does not contribute to the X:A ratio in the germ line. *Development* **117**, 763–767.
- Steinmann-Zwicky M., Schmid H. and Nöthiger R. 1989 Cell autonomous and inductive signals can determine the sex of the germ line of *Drosophila* by regulating the gene *Sxl. Cell* 57, 157–166.
- Steinmann-Zwicky M., Amrein H. and Nöthiger R. 1990 Genetic control of sex determination in *Drosophila*. Adv. Genet. 27, 189–237.
- Warn R. M., Magrath R. and Webb S. 1985 Distribution of F-actin during cleavage of *Drosophila* syncitial blastoderm. J. Cell. Biol. 98, 156–162.
- Wieschaus E., Audit C. and Masson M. 1981 A clonal analysis of the roles of somatic cells and germ line during oogenesis in *Drosophila. Dev. Biol.* 88, 92–103.

Received 17 March 1998; revised version received 22 September 1999