Regulation of sex-specific RNA splicing at the *Drosophila doublesex* gene: *cis*-acting mutations in exon sequences alter sex-specific RNA splicing patterns

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Sex-specific alternative RNA splicing of the *doublesex* (*dsx*) pre-mRNA results in sex-specific polypeptides that regulate both male and female somatic sexual differentiation in *Drosophila melanogaster*. We have molecularly characterized a class of *dsx* mutations that act in *cis* to disrupt the regulation of *dsx* RNA processing, causing the *dsx* pre-mRNA to be spliced in the male-specific pattern regardless of the chromosomal sex of the fly. These *dsx* mutations are associated with rearrangements in the female-specific exon just 3' to the female-specific splice acceptor. The mutations do not affect the female-specific splice sites or intron that are identical to wild-type sequences. These results indicate that sequences in the female-specific exon are important for the regulation of sex-specific RNA splicing, perhaps by acting as sites of interaction with *trans*-acting regulators.

Furthermore, the data suggest that female-specific regulation of *dsx* RNA processing occurs by promoting the usage of the female splice acceptor site, rather than by repressing the usage of the alternative male-specific splice acceptor.

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RNA splicing occurs through the interaction of *trans*-acting factors, such as the individual components of spliceosomes, with conserved *cis*-acting sequences found in pre-mRNAs. These sequences include the splice donor, the splice acceptor, and the branchpoint, all of which are required for correct splicing in both in vitro and in vivo systems (for review, see Green 1986; Padgett et al. 1986; Sharp et al. 1987a, b). In a growing number of genes, however, RNA splicing is complicated by the fact that a single pre-mRNA can be differentially spliced [Breitbart et al. 1987]. The usage of alternative RNA splice sites provides a mechanism by which a single gene can produce multiple, structurally related proteins whose expression can be regulated. Well-characterized examples of regulated splicing include the vertebrate calcitonin—cGRP gene [Amara et al. 1982; Sabate et al. 1985; Leff et al. 1987], the mammalian fibronectin gene [Kornbluth et al. 1985; Scharzbauer et al. 1987; Barone et al. 1989], the IgM heavy-chain gene [Alt et al. 1980; Early et al. 1980; Rogers et al. 1980; Galli et al. 1987, 1988; Peterson and Perry 1989], and the adenovirus E3 transcription unit [Nevins and Chen-Kiang 1981; Adami and Nevins 1988]. Alternative RNA processing can also be used as an 'on-off switch,' in which one splicing from encodes a protein product and an alternative form does not [Boggs et al. 1987; Bell et al. 1988; for review, see Bingham et al. 1988].

How the choice is made between differentially used splice sites is not well understood. A priori expectations are that *cis*-acting regulatory elements are required to confer differences between the alternatively used RNA splice sites to create asymmetry in their availability to the cellular splicing machinery. A number of *cis*-acting regions have been proposed to be involved in splice site choice, these appear to differ depending on the gene studied. For example, in the calcitonin—cGRP gene, differential poly(A) site selection and exon usage lead to alternative mature transcripts. This regulated RNA processing depends on sequences at the differentially used 3' splice acceptor but appears to be independent of sequences at the poly(A) sites [Leff et al. 1987; Adami and Nevins 1988]. This observation leads to the suggestion that *trans*-acting regulatory factors act to promote the usage of one splice acceptor over another which, in turn, determines the poly(A) site to be used. In comparison, two different models have been proposed for the regulation of differential RNA processing at the IgM gene, which also exhibits alternative 3' exon usage and polyadenylation. Galli et al. [1987, 1988] suggested that the regulation of IgM occurs at the level of poly(A) site

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choice which, in turn, determines the splicing pathway that is taken. An alternative model was proposed by Peterson and Perry (1989). They presented evidence that the IgM RNA splice pattern is determined by competition between the use of a nonconsensus IgM splice donor and a nearby poly(A) site. The choice between these mutually exclusive sites is dependent on cell-specific, trans-acting factors. Additional sites required in cis for splice site selection have been implicated in studies on other loci, including the branch site (Noble et al. 1987; Fu et al. 1988) and sequences within the alternatively spliced exons (Somasekhar and Mertz, 1985; Mardon et al. 1987; Barone et al. 1989; Hampson et al. 1989; Laski and Rubin 1989; Streuli and Saito 1989). This diversity of regions involved in regulating splice choice suggests that multiple mechanisms exist for controlling RNA processing.

A complete understanding of how RNA processing is regulated will require the characterization of both cis- and trans-acting factors that regulate RNA processing site choice. One of the best systems for studies of this kind is the regulatory pathway that controls somatic sexual development in Drosophila. Extensive genetic and molecular studies have demonstrated that the sex-specific regulation of this developmental pathway is controlled primarily, if not solely, by sex-specific RNA splicing (for review, see Baker 1989; Hodgkins 1989). Particularly advantageous is that at least some of the loci required in trans for the sex-specific splicing of these genes have been identified.

In Drosophila, the choice between male or female somatic development is determined by the assessment of the X/A ratio, the number of X chromosomes relative to the number of sets of autosomes. The X/A ratio determines somatic sexual differentiation by ultimately controlling the expression of the bifunctional doublesex (dsx) gene (Baker and Ridge 1980). In chromosomally female flies (X/A = 1 : 1), a hierarchy of regulatory genes act in response to the X/A ratio to cause dsx to be expressed in a female-specific manner. The dsx female product represses female differentiation. In contrast, in male flies (X/A = 1 : 2) dsx is expressed in a male-specific manner and the dsx male product acts to repress female differentiation. In this regard dsx is unique among the loci involved in somatic sexual differentiation in that it is the only gene whose active function is required to regulate the development of both sexes (Baker and Ridge 1980). Loss-of-function mutations of dsx cause intersexual development in both chromosomally male and female flies.

The basis of the bifunctionality of the dsx gene derives from sex-specific RNA processing (Burris and Baker 1989). The dsx gene produces both male- and female-specific mRNAs that are derived from a common pre-mRNA and have the same first three 5’ exons [Fig. 1]. They differ in their 3’ exons as a consequence of the use of sex-specific splice acceptor and poly(A) sites. One female RNA of 3.5 kb and two male-specific transcripts of 2.9 and 3.9 kb are detected by Northern blot analysis. The two male RNAs arise from differential poly(A) site usage in the last male-specific, noncoding exon. This male- and female-specific alternative RNA processing results in sex-specific polypeptides that differ in their carboxy-terminal domains.

Male-specific RNA splicing and polyadenylation of the dsx pre-mRNA is thought to be thedefault state of dsx RNA processing, because it occurs in the absence of activity of the genes known to regulate sexual differentiation (Baker et al. 1987; Nagoshi et al. 1988). In contrast, female-specific RNA processing at dsx is dependent on the activity of the transformer (tra) and transformer-2 (tra-2) genes [Fig. 2], as loss-of-function mutations of tra or tra-2 in chromosomally female flies results in the male dsx RNA splicing pattern [Nagoshi et al. 1988]. Although it has not been established that the tra and tra-2 products interact directly with the dsx pre-mRNA to impose female-specific RNA splicing, a direct interaction is suggested by the fact that the predicted tra-2 polypeptide contains a 90-amino-acid domain found in many proteins known to bind RNA (Amrein et al. 1988; Goralski et al. 1989). In addition, the predicted tra-2 (Goralski et al. 1989; Mattox et al. in prep) and tra (Belote et al. 1989) proteins also contain stretches of arginine- and serine-rich regions that are characteristic of some proteins that are involved in, or that may regulate, RNA processing (Chou et al. 1987; Bingham et al. 1988; Query et al. 1989).

We are interested in identifying and characterizing cis-acting dsx sequences that are important for sex-specific RNA processing. These are potential sites of interaction for trans-acting regulatory factors such as the tra and tra-2 products. In this paper we describe a class of cis-acting mutations at dsx that alter the dsx sex-specific RNA splicing pattern. We describe the phenotype, location, and structure of these lesions and discuss their potential role in the regulation of dsx RNA processing.

Figure 1. Model for the regulation of dsx. The dsx gene displays regulated RNA splicing and polyadenylation, which results in sex-specific polypeptides (Burris and Baker 1989). In chromosomally female flies, where the ratio of the number of X chromosomes to sets of autosomes (X/A) is 1 : 1, the tra and tra-2 genes act to cause female-specific RNA processing of the dsx gene (Nagoshi et al. 1988). In chromosomally male flies (X/A = 1 : 2), the tra gene is inactive. In the absence of female-specific regulation, the dsx gene is expressed in the default male mode. RNAs are not drawn to scale. [Open boxes] Exons; [boxes outlined by broken lines] unspliced exons; [shaded boxes] open reading frames; [AAA] polyadenylation sites; [diagonal lines] splice pattern; [broken line] introns.
Results

dsx dominant mutations

Genetic studies have identified a group of four dominant mutations at dsx that are distinct from the recessive loss-of-function dsx mutations in their phenotypic effects (Duncan and Kaufman 1975; Baker and Ridge 1980; Nothiger et al. 1980, 1987). The dsx dominant mutations are designated dsx\textsuperscript{O}, dsx\textsuperscript{M}, dsx\textsuperscript{T}, and dsx\textsuperscript{S} and are independent in origin but have nearly identical mutant phenotypes (summarized in Table 1). The dsx dominant mutations have two effects on dsx gene expression: (1) They eliminate the female-specific dsx function, and (2) they cause constitutive expression of the male dsx function. Thus, chromosomally female (XX) flies hemi-
zygous for the dsx dominant mutations [dsx\textsuperscript{dom}/Df(dsx)] develop somatically as males. However, when the dsx dominant alleles are heterozygous with a wild-type dsx allele, they transform XX flies into intersexes that are indistinguishable from those derived from homozygous recessive dsx-null mutations [see below]. In contrast, no mutant phenotype is found in chromosomally male (XY) flies carrying a dsx dominant mutation, either hemi-
zygous or over a wild-type dsx allele. This is consistent with the dominant dsx alleles expressing the male-spe-
cific dsx function that is normally active in male flies.

Table 1. dsx dominant mutant phenotypes

| Somatic phenotype | dsx function |
|-------------------|--------------|
| XX\textsuperscript{+}\textsuperscript{+} | male | + | - |
| Df(dsx)\textsuperscript{b} | male | + | - |
| XX\textsuperscript{+-} | intersex | + | + |
| XX\textsuperscript{-}\textsuperscript{-} | intersex | - | - |
| XY\textsuperscript{Om} | male | + | - |
| Df(dsx)\textsuperscript{b} | male | + | - |
| XY\textsuperscript{+}\textsuperscript{+} | intersex | + | - |
| XY\textsuperscript{-}\textsuperscript{-} | intersex | - | - |

The somatic phenotypes of the dsx dominant mutations are des-
cribed. The presence (+) or absence (-) of dsx function is ex-
trapolated from the sexual phenotype of the fly.

\textsuperscript{a}All four dominant mutations gave similar phenotypes.

\textsuperscript{b}Df(dsx) is dsx\textsuperscript{M+Rls} [B. Baker et al., in prep.], a deletion for the entire dsx locus.

To determine how the dsx dominant mutations disrupt dsx expression, we first examined the effects of these mutations at the transcript level. In particular, because previous studies demonstrated that sex-specific regulation at dsx is mediated through the control of RNA pro-
cessing, we examined whether the dsx dominant mutations disrupt the regulation of dsx RNA splicing.

The dsx dominant mutations alter the types of dsx transcripts produced in chromosomally female flies. This is illustrated by Northern blot analysis using single-stranded DNA probes specific for either male [Fig. 2A] or female dsx transcripts [Fig. 2B]. The wild-type pattern is shown in lane 9 for males and lane 10 for fe-
male. Lanes 1–4 contain RNA from chromosomally fe-
male flies that carry a dsx dominant allele over a dele-
tion of the dsx locus. Each of the four dominant alleles produces the male-specific dsx transcripts [Fig. 2A] and not the female-specific dsx transcript [Fig. 2B], even though these flies are chromosomally female. In chro-
mosomally female flies heterozygous for a dsx dominant and a dsx\textsuperscript{+} allele [lanes 5–8], both male- and female-spe-
cific transcripts are present—the female transcript from the dsx\textsuperscript{+} allele and the male transcript from the dsx dominant chromosome.

These results demonstrate that the dsx dominant mutations are cis-acting lesions that not only eliminate the
female transcript but also cause the male dsx transcript to be expressed in both chromosomally female and male flies. This indicates that the two phenotypes displayed by the dsx dominant mutations, the absence of female dsx function, and the constitutive expression of male dsx function are the consequences of an abnormal pattern of dsx RNA processing. The ectopic expression of the male dsx transcript in chromosomally female flies argues against the dominant lesions affecting the stability or accumulation of the female-specific dsx RNA. Rather, it appears that the pre-mRNA from the dsx dominant alleles is incapable of female-specific splicing and polyadenylation, instead displaying the default [male] RNA processing pattern.

The dsx RNA splicing pattern in flies hemizygous for a dsx dominant allele is similar to that found in flies lacking tra or tra-2 activity. In both cases, the absence of the female-specific dsx RNA is associated with the presence of the male-specific dsx transcript. A simple explanation for these results is that the dsx dominant mutations are cis-acting lesions that prevent tra and tra-2, or factors regulated by these genes, from interacting with the dsx pre-mRNA. This would prevent female-specific regulation of dsx RNA processing, thereby causing constitutive usage of the default male splice acceptor site.

Mapping of the dsx dominant mutations

Because the dsx dominant lesions act in cis to disrupt sex-specific RNA processing, it is likely that they are located in regions that are important for sex-specific regulation. The mapping of these lesions should provide insight into the mechanism by which dsx RNA processing is controlled. Two topics, in particular, can be addressed. First, because the dsx pre-mRNA undergoes both sex-specific splicing and polyadenylation, the regulation of dsx RNA processing could be dictated by the choice of splice acceptor, poly[A] site, or a combination of the two. The clustering of the dsx dominant lesions in the vicinity of one of these RNA processing sites would argue for its importance in the regulation of dsx sex-specific expression. Second, the regulation of dsx RNA processing by trans-acting factors, such as tra and tra-2, could occur by either positive or negative mechanisms (Fig. 3). In chromosomally female flies, the wild-type tra and tra-2 products could have a negative regulatory role by acting at the male splice acceptor [or poly[A] site] to prevent male-specific RNA processing from occurring. This would then allow the alternative female splice acceptor to be used. On the other hand, tra and tra-2 could act positively to promote the usage of the female-specific processing sites instead of the default male-specific sites. The location of the dsx dominant lesions relative to the male- and female-specific RNA processing sites could help distinguish between these possibilities.

The initial localization of the dsx dominant mutations was done by restriction fragment length polymorphism (RFLP) analysis, using a combination of both Southern blotting of genomic DNAs and the DNA cloning and characterization of each of the four dsx dominant alleles (Baker and Wolfrner 1988; this paper). All four alleles displayed restriction pattern alterations from the wild-type map within an ~3-kb interval containing the female-specific splice acceptor and polyadenylation sites. Across the 40-kb dsx locus, this was the only region where all four dominant alleles differed in their restriction pattern from the wild-type Canton S pattern. The correlation in four independent mutations between the location of a rearrangement in a discrete region of the dsx gene and the dsx dominant mutant phenotype provides strong support for the inference that these rearrangements are the mutations.

Sequence analysis of the mutant DNAs precisely localized the structural lesions associated with each of the dominant mutations. All four dominant alleles are associated with rearrangements with breakpoints in the vicinity of the female-specific splice acceptor (Fig. 4). The dsxD, dsxE, and dsxH alleles are each associated with an insertion of middle repetitive elements that lie 48, 127, and 175 bp, respectively, 3' of the female-specific splice acceptor site and within the female-specific exon. The dsxH allele carries a 448-bp deletion associated with the addition of five bases: CCATA. The 5' breakpoint of the deletion is located 108 bp, 3' of the female-specific splice acceptor. These rearrangements are completely localized within the female-specific exon. In contrast, the female-specific splice sites and the entire female-specific intron were found to be identical to the wild-type DNA sequence for all four dominant mutations.

Unlike the close proximity of these rearrangements to the female splice acceptor, all are located >600 bp upstream from the female canonical AATAAA polyadenylation signal. The most distant rearrangement, the dsxO
this repeat may be involved in the interaction of the dsx pre-mRNA with trans-acting regulators is currently being tested.

Characterization of the dsx dominant-associated insertions

The insertions associated with dsx⁰, dsx⁴, and dsx⁷ are described in Table 2. Both the dsx⁰ and dsx⁷ insertions are associated with a 20- to 25-residue stretch of poly[A] at one of their ends, a characteristic of certain families of middle repetitive retroposons in Drosophila melanogaster [Di Nocera et al. 1983; Pittler and Davis 1987; Di Nocera 1988]. The dsx⁴ element shares substantial sequence homology with one such family of repeats, the F elements [Table 2B; Di Nocera et al. 1983], in the region immediately adjacent to the poly[A] stretch. However, the dsx⁴ element may represent a different subgroup of this family because no homology could be found with that of the consensus F element in the sequence of the end opposite the poly(A) stretch. No sequence homologies could be found for the dsx⁷ element with the F family or with the other sequences contained in the National Institutes of Health [NIH] GenBank [release 58.0] and EMBL Nucleotide Sequence Data Library [release 17.0]. Thus, it may represent a new family of RNA-mediated transposable elements. The dsx⁰ insertion was found to have substantial sequence similarity with the roo transposable element [Table 2B; Scher rer et al. 1982, Garfinkle et al. 1983]. It is 7.8 kb in size and is present in ~100 copies per genome, as estimated from genomic Southern blots.

Male and female dsx products are antagonistic

On the basis of genetic studies that indicate that the male- and female-specific dsx products act to repress female and male sexual differentiation, respectively, the simultaneous expression of both products should repress sexual differentiation altogether. The expected phenotype of this situation is difficult to predict; one might expect the absence of sexually dimorphic tissues or that these tissues would develop but be sexually undifferentiated. Instead, the dsx dominant mutation over dsx⁺ genotype results in an intersexual phenotype that is indistinguishable from that resulting from dsx null mutations [Table 1; Baker and Ridge 1980, Nothiger et al. 1987]. This suggests that the male and female dsx products are mutually antagonistic; their simultaneous expression in a single cell is functionally equivalent to the absence of dsx products. From these observations, each of the dsx sex-specific products can be seen to have two activities: (1) When only one of the products is expressed, it represses differentiation of the opposite sex; and (2) when both dsx products are expressed, they inactive each other. The reason for this mutual antagonism is not known.

The data in Figure 2 indicate that the antagonism between the activities of the dsx male and female products does not occur at the level of RNA synthesis or pro-
Table 2. Sequence analysis of the \( dsx \) dominant lesions

| Allele | Proximal end | Distal end | Repeated sequence at site of insertion | Insert size (kb) |
|--------|--------------|------------|---------------------------------------|-----------------|
| \( dsx^D \) | ACAAGTGTGAATCTTTGGTTTCAATATAATTAAAGAACACTTT | TCTAAATCCCATATAAGAAGA | GCTTC | 7.7 |
| \( dsx^M \) | \( \{A\}_{13}^{A} \) TGAAGGAAATTAAATTAATATCAGA | TCCATT | ATAA | 3.9 |
| \( dsx^F \) | GCCGATTCTCTGTAGTCCATATCTCTGC | ACAAGTGTGAATCTTTGGTTTCAATATAATTAAAGAACACTTT | TTTTGAATTTTGCTCTATA | TTTGCC | 4.5 |
| \( dsx^M \) | \( \{A\}_{13}^{A} \) TGAAGGAAATTAAATTAATATCAGA | TCCATT | ATAA | 3.9 |

Sequence comparison of the \( dsx^M \) and \( dsx^D \) insertions with known transposable elements

\( \text{F element}^a \)

\( ^b \)From Scherer et al. (1982).

\( ^c \)From Di Nocera et al. (1983).

In chromosomally female flies carrying both a \( dsx \) dominant mutation with a wild-type \( dsx \) allele, both male- and female-specific \( dsx \) transcripts are expressed [Fig. 2, lanes 5–8]; the male product from the \( dsx \) dominant allele and the female product from the \( dsx^+ \) gene. This indicates that the presence of the female \( dsx \) product does not repress the production of the male \( dsx \) RNA nor does the male \( dsx \) product inhibit female-specific \( dsx \) RNA processing.

Discussion

In this study we examined a unique class of \( dsx \) mutations, the \( dsx \) dominant alleles, which alter two aspects of \( dsx \) gene expression: (1) They eliminate the female-specific \( dsx \) function, and (2) they cause constitutive expression of the male-specific \( dsx \) function. Both of these mutant phenotypes are the consequences of alterations in the pattern of \( dsx \) RNA processing such that only male-specific splicing and polyadenylation occurs in both chromosomally male and female flies. To explain these results, we propose the following mechanism for the action of the \( dsx \) dominant mutations and the potential role of the \( tra \) and \( tra-2 \) genes.

The male-specific splicing pattern is a default state for \( dsx \) RNA processing, it occurs in the absence of sex-specific regulatory functions requiring only the general housekeeping splicing machinery of the cell [Baker et al. 1987; Nagoshi et al. 1988]. In the absence of sex-specific regulation, the male splice acceptor is preferentially used over the female acceptor, perhaps reflecting the closer similarity of the male splice acceptor to the \( Drosophila \) consensus splice acceptor sequence as compared to that shown by the female splice acceptor [Burtis and Baker 1989]. We propose that female-specific trans-acting regulatory factors, particularly the \( tra \) and \( tra-2 \) gene products, act to overcome this preferential usage of the male acceptor, thereby allowing the female splice to occur in chromosomally female flies. The \( dsx \) dominant mutations prevent this regulation from occurring, perhaps by disrupting regions in the \( dsx \) RNA required for binding to the \( tra \) and \( tra-2 \) products. Without the intervention of these trans-acting factors, the \( dsx \) primary transcript is processed in the default pattern, resulting in male-specific splicing and polyadenylation. This mechanism explains how a single lesion in the \( dsx \) gene, the \( dsx \) dominant mutations, can both prevent female-specific RNA processing and cause constitutive production of the male \( dsx \) transcripts.

On the basis of the above model, there are two possible ways by which the regulation of female-specific splicing could occur. Trans-acting regulators, such as \( tra \) and \( tra-2 \), could act in a positive fashion to promote the usage of the nonconsensus female-specific splice site. Alternatively, the female-specific regulators could act negatively by preventing the usage of the default male splice acceptor. The proximity of the dominant lesions to the female splice acceptor is most consistent with \( tra \) and \( tra-2 \) promoting the usage of the female splice site. Furthermore, the locations of the dominant lesions implicate exon sequences within 200 bp 3' of the female-specific splice acceptor to be required for the regulation of female-specific splicing.

There is precedent for the involvement of exon sequences in splice acceptor site selection [Somasekhar and Mertz 1985; Reed and Maniatis 1986]. Reed and Maniatis (1986) demonstrated with in vitro studies that if two identical splice acceptors are placed in tandem array downstream from a splice donor, the nearest splice acceptor is preferentially used. However, this preference can be shifted to the more distant splice acceptor by altering exon sequences immediately 3' of the first splice acceptor. In the human fibronectin gene, sequences within a differentially spliced exon were found to be required in vivo for regulated RNA splicing [Barone et al. 1989]. The deletion or reversal of orientation of an 81-bp
region in the middle of this exon prevented the usage of the adjacent splice acceptor site. A similar phenomenon was described for the human leukocyte common antigen (LCA) RNA in in vivo experiments [Streuli and Saito 1989]. The LCA pre-mRNA undergoes cell-type-specific usage of splice acceptor sites, the regulation of which is dependent on the physical integrity of several discrete regions in the differentially spliced exon.

Reed and Maniatis [1986] suggested that exon sequences could participate directly in splice site recognition by determining the affinity of adjacent splice sites for splicing components. This could occur through RNA secondary structure, perhaps by making the splice site conformationally more accessible, or alternatively, the exon sequences themselves might bind directly to factors involved in splice site selection. The latter possibility seems likely in the current case, because of the finding that two genes that may directly control sex-specific RNA processing at dsx, tra (Belote et al. 1989), and tra-2 (Amrein et al. 1988; Goralski et al. 1989) encode for polypeptides with structural similarities to known RNA-binding proteins.

An alternative possibility is that the spacing between the sex-specific splice acceptors and polyadenylation sites is critical for sex-specific regulation. There could be, for example, a functional interaction between female-specific splicing and polyadenylation such that variations in the spacing of the splice acceptor from the polyadenylation site would have dramatic effects on splice site selection. Alternatively, there may be competition between the usage of the male and female splice acceptors (or polyadenylation sites) whose resolution is, in part, dependent on the physical distance between the sites. A potential example of this type of mechanism is the IgM heavy-chain gene which, like dsx, displays both alternative splice acceptor and alternative poly[A] site selection [Alt et al. 1980; Early et al. 1980; Rogers et al. 1980]. Poly[A] site selection was reported to be dependent on the spacing between the alternative poly[A] sites (Galli et al. 1987, 1988). Another study indicated that changes in the relative locations of the differential splice donor to the poly[A] sites can affect RNA splice site selection [Peterson and Perry 1986, 1989]. The occurrence of a similar mechanism for the regulation of dsx seems unlikely in view of the fact that either insertions of as much as 7.8 kb [dsx<sup>+</sup>] or a deletion of 0.5 kb [dsx<sup>-</sup>] within the region that separates the female splice acceptor from both the male splice acceptor and female polyadenylation site result in identical patterns of dsx pre-mRNA splicing.

In summary, we have identified cis-acting sequences in the dsx gene that have profound effects on the regulation of sex-specific splicing of the dsx RNA. These are likely to be sites of interaction with the trans-acting regulators of dsx RNA processing, two likely candidates for such factors are the products of the tra and tra-2 genes. The delineation of how these cis- and trans-acting factors interact to regulate sex-specific splicing at dsx should have general relevance toward understanding how RNA processing is regulated.

Methods

Fly strains

The genetic phenotypes of the dsx<sup>T</sup> [Duncan and Kaufman 1975] and dsx<sup>M</sup> (Baker and Ridge 1980, Nothiger et al. 1980) alleles have been described previously. The dsx<sup>T</sup> allele was isolated in W. Gehring’s laboratory in Basel. All three are spontaneous in origin. The dsx<sup>M</sup> allele was induced by X-ray mutagenesis in an inbred background for which the parental chromosome is available [K. Brandt-Rosquist, pers. comm., as cited in Baker and Wollner 1988]. The deletion, dsx<sup>M<sup>−</sup>415</sup>, removes the entire dsx gene and is described in B. Baker et al. [in prep.]. Flies were raised on a standard cornmeal, molasses, yeast, agar medium, containing propionic acid as a mold inhibitor and supplemented with live yeast.

RNA preparation

RNAs for Northern blot analyses were obtained as follows: Adult flies (10–50) were homogenized in a Dounce homogenizer in 10 ml of 1 : 1 homogenization buffer/phenol [homogenization buffer: 0.15 M NaCl, 1.5 mM MgCl<sub>2</sub>, 10 mM Tris-HCl (pH 8.0), 0.5% SDS]. The supernatant was extracted twice with 1 : 1 phenol/chloroform and ethanol-precipitated. The nucleic acid precipitate was resuspended in 10 mM Tris-HCl (pH 8.0) and 1 mM EDTA and the concentration determined by OD<sub>260</sub>.

RNA blotting and hybridization

Northern analysis was done as described in Nagoshi et al. [1988]. Total RNA (10–20 μg per lane) was loaded in a formaldehyde gel and blotted to Nytran [Schleicher & Schuell], as described in Maniatis et al. [1982]. Single-stranded DNA probes were derived from dsx genomic and cDNA sequences subcloned into the M13 vector, using primed synthesis from a 17-mer M13 primer. Hybridizations were done in 5 x SSPE (or 5 x SSC), 5 x Denhardt’s solution, 50 μg/ml salmon sperm DNA, and 0.2% SDS at 42°C.

DNA sequencing

Genomic DNA fragments were subcloned into either M13 [Yanisch-Perron et al. 1985] or Bluescript (Strategene) vectors. Single- and double-stranded templates were prepared by standard methods and sequenced by the method of Sanger et al. [1977]. Computer resources for sequences were provided by the Bionet National Computer Resources for Molecular Biology, which is funded by the Biomedical Research Technology Program, Division of Research Resources, National Institutes of Health.

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Nagoshi and Baker

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R N Nagoshi and B S Baker

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